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(54) Title: IMMUNOMODULATORY PEPTIDES

### (57) Abstract

A purified preparation of a peptide consisting essentially of an amino acid sequence identical to that of a segment of a naturally-occurring human protein, said segment being of 10 to 30 residues in length, inclusive, wherein said peptide binds to a human major histocompatibility complex (MHC) class II allotype.

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## IMMUNOMODULATORY PEPTIDES

This application is a continuation-in-part of copending USSN 07/925,460, filed August 11, 1992. The invention was made in the course of research funded in part by the U.S. Government under NIH Grant 5R35-CA47554; the U.S. Government therefore has certain rights in the invention.

The field of the invention is major 10 histocompatibility complex (MHC) antigens.

# Background of the Invention

Major histocompatibility complex (MHC) class II antigens are cell surface receptors that orchestrate all specific immune responses in vertebrates. Humans possess three distinct MHC class II isotypes: DR, for which approximately 70 different allotypes are known; DQ, for which 33 different allotypes are known; and DP, for which 47 different allotypes are known. Each individual bears two to four-DR alleles, two DQ alleles, and two DP 20 alleles.

MHC receptors (both class I and class II)

participate in the obligate first step of immune

recognition by binding small protein fragments (peptides)

derived from pathogens or other non-host sources, and

25 presenting these peptides to the regulatory cells (T

cells) of the immune system. In the absence of MHC

presentation, T cells are incapable of recognizing

pathogenic material. Cells that express MHC class II

receptors are termed antigen presenting cells (APC).

30 APCs ingest pathogenic organisms and other foreign

materials by enveloping them in endosomic vesicles, then

subjecting them to enzymatic and chemical degradation.

Foreign proteins which are ingested by APCs are partially

degraded or "processed" to yield a mixture of peptides,

35 some of which are bound by MHC class II molecules that

are en route to the surface. Once on the cell surface, MHC-bound peptides are available for T cell recognition.

MHC class II antigens are expressed on the surface of APCs as a trimolecular complex composed of an  $\alpha$  chain, 5 a  $\beta$  chain, and a processed peptide. Like most polypeptides that are expressed on the cell surface, both  $\alpha$  and  $\beta$  chains contain short signal sequences at their NH2 termini which target them to the endoplasmic reticulum (ER). Within the ER the class II  $\alpha/\beta$  chain 10 complex associates with an additional protein termed the invariant chain (Ii). Association with Ii is proposed to block the premature acquisition of peptides (by blocking the peptide binding cleft of the MHC heterodimer), promote stable  $\alpha/\beta$  interaction, and direct subsequent 15 intracellular trafficking of the complex to endosomal vesicles. In the endosomes, Ii is removed by a process involving proteolysis; this exposes the peptide binding cleft, thus allowing peptides present in the endosome to bind to the MHC molecule. The class II/ peptide complex 20 is transported from the endosomes to the cell surface where it becomes accessible to T-cell recognition and subsequent activation of immune responses. Class II MHC molecules bind not only to peptides derived from exogenous (ingested) proteins, but also to those produced 25 by degradation of endogenous (self) proteins. The amount of each species of peptide which binds class II is determined by its local concentration and its relative binding affinity for the given class II binding groove, with the various allotypes displaying different peptide-30 binding specificities.

Early during fetal development, the mammalian immune system is "tolerized", or taught not to react, to self-peptides. The stability and maintenance of this system is critical for ensuring that an animal does not generate an immune response against self. A breakdown of

this system gives rise to autoimmune conditions such as diabetes, rheumatoid arthritis and multiple sclerosis. Current technologies intended to manipulate the immune system into reestablishing proper nonresponsiveness include protocols involving the intravenous delivery of synthetic, high affinity binding peptides as blocking peptides.

Vaccination can generate protective immunity against a pathogenic organism by stimulating an antibody10 mediated and/or a T cell-mediated response. Most of the current vaccination strategies still use relatively crude preparations, such as attenuated or inactivated viruses. These vaccines often generate both antibody- and cell-mediated immunity, and do not allow one to modulate the type of immune response generated. Moreover, in many diseases the generation of the wrong type of response can result in an exacerbated disease state.

# Summary of the Invention

In the work disclosed herein, naturally processed 20 peptides bound to six of the some 70 known human MHC class II DR allotypes (HLA-DR1, HLA-DR2, HLA-DR3, HLA-DR4, HLA-DR7, and HLA-DR8) have been characterized. These peptides were found to be predominantly derived from self proteins rather than foreign proteins. Several 25 self peptide families have been identified with the unexpected property of degenerate binding: that is, a given self-peptide will bind to a number of HLA-DR allotypes. This observation runs counter to the widelyaccepted view of MHC class II function, which dictates 30 that each allotype binds a different set of peptides. Furthermore, many if not all of the self-peptides disclosed herein bind to the class II molecules with relatively high affinity. These three characteristics--(1) self rather than foreign, (2) degeneracy, and (3)

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high affinity binding--suggest a novel means for therapeutic intervention in disease conditions characterized by autoreactivity, such as Type I diabetes, rheumatoid arthritis, and multiple sclerosis. In addition, such therapy could be used to reduce transplant rejection.

In the therapeutic methods of the invention, short peptides modelled on the high-affinity immunomodulating self peptides of the invention (which preferably are 10 nonallelically restricted) are introduced into the APCs of a patient. Tissue typing to determine the particular class II alleles expressed by the patient may be unnecessary, as the peptides of the invention are bound by multiple class II isotypes. It may be useful to 15 employ a "cocktail" of peptides, where complete degeneracy is lacking for individual peptides, i.e., where peptides binds to fewer than all allotypes; the cocktail provides overlapping binding specificity. Once in the APC, a peptide binds to the class II molecules 20 with high affinity, thereby blocking the binding of immunogenic peptides which are responsible for the immune reaction characteristic of the disease condition. Because the blocking peptides of the invention are self peptides with the exact carboxy and amino termini 25 tolerized during ontogeny, they are immunologically inert and will not induce an immune response which may complicate treatment using non-self blocking peptides.

The peptides of the invention may be introduced into APCs directly, e.g., by intravenous injection of a 30 solution containing one or more of the peptides.

Alternatively, the APCs may be provided with a means of synthesizing large quantities of the blocking peptides intracellularly. Recombinant genes that encode ER and/or endosomal targeting signals fused to blocking peptide 35 sequences are linked to appropriate expression control

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sequences and introduced into APCs. Once in the cell, these genes direct the expression of the hybrid peptides. Peptides targeted to the ER will bind class II  $\alpha$  and  $\beta$ chains as they are translated and assembled into 5 heterodimers. The presence of high affinity binding peptides within the ER will prevent association of the  $\alpha/\beta$  complex with invariant chain, and thus interfere with intracellular trafficking. The class II molecule/ blocking peptide complex may subsequently be expressed on 10 the cell surface, but would not elicit an immune response since T cells are tolerized to this complex early in development. The use of peptides tagged with ER retention signals may also prevent the peptide-complexed class II molecules from leaving the ER. Alternatively, 15 the recombinant peptide may be tagged with an endosomal targeting signal which directs it to the endosomal compartment after synthesis, thereby also skewing the ratio of endogenously-processed peptide to blocking peptide in the endosome and favoring binding of the high 20 affinity blocking peptide to any class II molecules which did not bind it in the ER. It may be advantageous, for any individual patient, to employ one or more ER-directed peptides in combination with one or more endosomedirected peptide, so that  $\alpha-\beta$  complexes which are not 25 filled in the ER with peptides of the invention are then blocked in the endocytic pathway. The end result again is cell surface expression of a non-immunogenic class II/peptide complex.

The use of a class II nonrestricted high affinity
30 binding peptide coupled to an intracellular delivery
system permits the specific down-regulation of class II
restricted immune responses without invoking the
pleiotropic adverse reactions associated with the current
pharmacological strategies. Successful application of
35 these technologies will constitute a significant advance

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towards the treatment of autoimmune disease and prevention of transplant rejection.

The intracellular delivery system of the invention can also be utilized in a novel method of vaccination of 5 an animal, e.g., a human patient or a commercially significant mammal such as a cow which is susceptible to diseases such as hoof and mouth disease. Such a system can be tailored to generate the type of immune response required in a given situation by adjustments in the 10 following: (a) peptide specificity for class I or class II MHC; (b) peptide/protein length and/or sequence, and (c) using specific tags for organelle targeting. The system of the invention ensures that peptides are produced only within cells, and are not present outside 15 the cells where they could stimulate antibody production by contact with B cells. This limits the immune response generated by such a vaccine to T cell-mediated immunity, thereby preventing either an inappropriate or potentially deleterious response as might be observed with standard 20 vaccines targeting the organisms which cause, for example, HIV, malaria, leprosy, and leishmaniasis. Furthermore, this exclusively T cell-mediated immune response can be class I or class II-based, or both, depending upon the length and character of the 25 immunogenic peptides: MHC class I molecules are known to bind preferentially to peptides 8 to 10 residues in length, while class II molecules bind with high affinity to peptides that range from 12 to 25 residues long.

Immunization and therapy according to the

invention can employ a purified preparation of a peptide
of the invention, i.e., a peptide which includes an amino
acid sequence identical to that of a segment of a
naturally-occurring human protein (i.e., a "self
protein"), such segment being of 10 to 30 residues in
length, wherein the peptide binds to a human MHC class II

allotype, and preferably binds to at least two distinct MHC class II allotypes (e.g., any of the approximately 70 known DR allotypes, approximately 47 known DP allotypes, or approximately 33 known DQ allotypes). The portion of 5 the peptide corresponding to the self protein segment is herein termed a "self peptide". By "purified preparation" is meant a preparation at least 50% (by weight) of the polypeptide constituents of which consists of the peptide of the invention. In preferred 10 embodiments, the peptide of the invention constitutes at least 60% (more preferably at least 80%) of the purified preparation. The naturally-occurring human protein is preferably HLA-A2 (as broadly defined below), HLA-A29, HLA-A30, HLA-B44, HLA-B51, HLA-Bw62, HLA-C, HLA-DP  $\beta$ -15 chain, HLA-DQ  $\alpha$ -chain, HLA-DQ  $\beta$ -chain, HLA-DQ3.2  $\beta$ -chain, HLA-DR α-chain, HLA-DR β-chain, HLA-DR4 β-chain, invariant chain (Ii), Ig kappa chain, Ig kappa chain C region, Ig heavy chain, Nat/Kt ATPase, potassium channel protein, sodium channel protein, calcium release channel 20 protein, complement C9, glucose-transport protein, CD35, CD45, CD75, vinculin, calgranulin B, kinase C ζ-chain, integrin  $\beta$ -4 gp150, hemoglobin, tubulin  $\alpha$ -1 chain, myosin  $\beta$ -heavy chain,  $\alpha$ -enolase, transferrin, transferrin receptor, fibronectin receptor α-chain, acetylcholine 25 receptor, interleukin-8 receptor, interferon α-receptor, interferon y-receptor, calcitonin receptor, LAM (lymphocyte activation marker) Blast-1, LAR (leukocyte antigen-related) protein, LIF (leukemia inhibitory factor) receptor, 4F2 cell-surface antigen (a cell-30 surface antigen involved in normal and neoplastic growth) heavy chain, cystatin SN, VLA-4 (a cell surface heterodimer in the integrin superfamily of adhesion receptors), PAI-1 (plasminogen activator inhibitor-1), IP-30 (interferon-y induced protein), ICAM-2,

35 carboxypeptidase E, thromboxane-A synthase, NADH-

cytochrome-b5 reductase, c-myc transforming protein, K-ras transforming protein, MET kinase-related transforming protein, interferon-induced guanylate-binding protein, mannose-binding protein, apolipoprotein B-100,

- 5 cathepsin C, cathepsin E, cathepsin S, Factor VIII, von Willebrand factor, metalloproteinase inhibitor 1 precursor, metalloproteinase inhibitor 2, plasminogen activator inhibitor-1, or heat shock cognate 71 kD protein; it may be an MHC class I or II antigen protein
- or any other human protein which occurs at the cell surface of APCs. The self peptide preferably conforms to the following motif: at a first reference position (I) at or within 12 residues of the amino terminal residue of the segment, a positively charged residue (i.e., Lys,
- 15 Arg, or His) or a large hydrophobic residue (i.e., Phe, Trp, Leu, Ile, Met, Tyr, or Pro; and at position I+5, a hydrogen bond donor residue (i.e., Tyr, Asn, Gln, Cys, Asp, Glu, Arg, Ser, Trp, or Thr). In addition, the peptide may also be characterized as having, at positions
- 20 I+9, I+1, and/or I-1, a hydrophobic residue (i.e., Phe, Trp, Leu, Ile, Met, Pro, Ala, Val, or Tyr) (+ denotes positions to the right, or toward the carboxy terminus, and denotes positions to the left, or toward the amino terminus.) A typical peptide of the invention will
- include a sequence corresponding to residues 31-40 (i.e., TQFVRFDSDA; SEQ ID NO: 149) or residues 106-115 (i.e., DWRFLRGYHQ; SEQ ID NO: 150) of HLA-A2, or residues 107-116 (i.e., RMATPLLMQA; SEQ ID NO: 151) of Ii, or a sequence essentially identical to any one of the sequences set forth in Tables 1-10 below.

The therapeutic and immunization methods of the invention can also employ a nucleic acid molecule (RNA or DNA) encoding a peptide of the invention, but encoding less than all of the entire sequence of the self protein.

35 The nucleic acid preferably encodes no substantial

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portion of the self protein other than the specified self peptide which binds to a MHC class II molecule, although it may optionally include a signal peptide or other trafficking sequence which was derived from the self 5 protein (or from another protein). A trafficking sequence is an amino acid sequence which functions to control intracellular trafficking (directed movement from organelle to organelle or to the cell surface) of a polypeptide to which it is attached. Such trafficking 10 sequences might traffic the polypeptide to ER, a lysosome, or an endosome, and include signal peptides (the amino terminal sequences which direct proteins into the ER during translation), ER retention peptides such as KDEL (SEQ ID NO: 152); and lysosome-targeting peptides 15 such as KFERQ (SEQ ID NO: 153), QREFK (SEQ ID NO: 154), and other pentapeptides having Q flanked on one side by four residues selected from K, R, D, E, F, I, V, and L. An example of a signal peptide that is useful in the invention is a signal peptide substantially identical to 20 that of an MHC subunit such as class II  $\alpha$  or  $\beta$ ; e.g., the signal peptide of MHC class II  $\alpha$  is contained in the sequence MAISGVPVLGFFIIAVLMSAQESWA (SEQ ID NO: 155). signal peptide encoded by the nucleic acid of the invention may include only a portion (e.g., at least ten 25 amino acid residues) of the specified 25 residue sequence, provided that portion is sufficient to cause trafficking of the polypeptide to the ER. In preferred embodiments, the nucleic acid of the invention encodes a second self peptide and a second trafficking sequence 30 (which may be identical to or different than the first self peptide and first trafficking sequence), and it may encode additional self peptides and trafficking sequences as well. In still another variation on this aspect of the invention, the self peptide sequence (or a plurality 35 of self peptide sequences arranged in tandem) is linked

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by a peptide bond to a substantially intact Ii polypeptide, which then carries the self peptide sequence along as it traffics the class II molecule from ER to endosome.

The nucleic acid of the invention may also contain expression control sequences (defined as transcription and translation start signals, promoters, and enhancers which permit and/or optimize expression of the coding sequence with which they are associated) and/or genomic nucleic acid of a phage or a virus, such as an attenuated or non-replicative, non-virulent form of vaccinia virus, adenovirus, Epstein-Barr virus, or a retrovirus.

The peptides and nucleic acids of the invention may be prepared for therapeutic use by suspending them

15 directly in a pharmaceutically acceptable carrier, or by encapsulating them in liposomes, amune-stimulating complexes (ISCOMS), or the like. Such preparations are useful for inhibiting an immune response in a human patient, by contacting a plurality of the patient's APCs with the therapeutic preparation and thereby introducing the peptide or nucleic acid into the APCs.

Also within the invention is a cell (e.g., a tissue culture cell or a cell, such as a B cell or APC, within a human) containing the nucleic acid molecule of the invention. A cultured cell containing the nucleic acid of the invention may be used to manufacture the peptide of the invention, in a method which involves culturing the cell under conditions permitting expression of the peptide from the nucleic acid molecule.

Disclosed herein is a method of identifying a nonallelically restricted immunomodulating peptide, which method includes the steps of:

(a) fractionating a mixture of peptides eluted from a first MHC class II allotype;

- (b) identifying a self peptide from this mixture; and
- (c) testing whether the self peptide binds to a second MHC class II allotype, such binding being an5 indication that the self peptide is a nonallelically restricted immunomodulating peptide.

In further embodiments, the invention includes a method of identifying a potential immunomodulating peptide, in a method including the steps of:

- 10 (a) providing a cell expressing MHC class II
   molecules on its surface;
  - (b) introducing into the cell a nucleic acid encoding a candidate peptide; and
- (c) determining whether the proportion of
  15 class II molecules which are bound to the candidate
  peptide is increased in the presence of the nucleic acid
  compared to the proportion bound in the absence of the
  nucleic acid, such an increase being an indication that
  the candidate peptide is a potential immunomodulating
  20 peptide.

Also within the invention is a method of identifying a potential immunomodulating peptide, which method includes the steps of:

- (a) providing a cell expressing MHC class II25 molecules on its surface;
  - (b) introducing into the cell a nucleic acidencoding a candidate peptide; and
- (c) determining whether the level of MHC class II molecules on the surface of the cell is decreased in the presence of the nucleic acid compared to the level of MHC class II molecules in the absence of the nucleic acid, such a decrease being an indication that the candidate peptide is a potential immunomodulating peptide.

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Also included in the invention is a method of identifying a nonallelically restricted immunostimulating peptide, which method includes the steps of:

- (a) providing a cell bearing a first MHC class I or class II allotype, such cell being infected with a pathogen (e.g., an infective agent which causes human or animal disease, such as human immunodeficiency virus (HIV), hepatitis B virus, measles virus, rubella virus, influenza virus, rabies virus, Corynebacterium diphtheriae, Bordetella pertussis, Plasmodium spp., Schistosoma spp., Leishmania spp., Trypanasoma spp., or Mycobacterium lepre);
  - (b) eluting a mixture of peptides bound to the cell's first MHC allotype;
- 15 (c) identifying a candidate peptide from the mixture, such candidate peptide being a fragment of a protein from the pathogen; and
- (d) testing whether the candidate peptide binds to a second MHC allotype, such binding being an 20 indication that the candidate peptide is a nonallelically restricted immunostimulating peptide. A nucleic acid encoding such an immunogenic fragment of a protein of a pathogen can be used in a method of inducing an immune response in a human patient, which method involves introducing the nucleic acid into an APC of the patient.

The therapeutic methods of the invention solve certain problems associated with prior art methods involving intravenous injection of synthetic peptides:

(1) because of allelic specificity, a peptide capable of binding with high affinity to all, or even most, of the different class II allotypes expressed within the general population had not previously been identified; (2) the half-lives of peptides delivered intravenously are generally very low, necessitating repeated administration

with the associated high level of inconvenience and cost;
(3) this type of delivery approach requires that the
blocking peptide displace the naturally-occurring peptide
occupying the binding cleft of a class II molecule while
the latter is on the cell surface, which is now believed
to be a very inefficient process; and (4) if the blocking
peptide utilized is itself immunogenic, it may promote
deleterious immune responses in some patients.

Other features and advantages of the invention
10 will be apparent from the following detailed description,
and from the claims.

## <u>Detailed Description</u>

The drawings are first briefly described.

## Drawings

Figs. 1A-1F are chromatographic analyses of the peptide pools extracted from papain digested HLA-DR1, DR2, DR3, DR4, DR7, and DR8, respectively, illustrating the peptide repertoire of each HLA-DR as detected by UV absorbance. The UV absorbance for both 210 nm and 277 nm is shown at a full scale absorbance of 500 mAU with a retention window between 16 minutes and 90 minutes (each mark represents 2 minutes).

Fig. 2 is a representative mass spectrometric analysis of the size distribution of isolated HLA-DR1 bound peptides. The determined peptide masses in groups of 100 mass units were plotted against the number of isolated peptides identified by mass spectrometry. Peptide length was calculated by dividing the experimental mass by an average amino acid mass of 118 daltons.

Fig. 3A is a representation of a minigene of the invention (SEQ ID NO: 147), in which the HLA-DRα chain leader peptide is linked to the amino terminus of a 15-residue blocking peptide fragment of human invariant 5 chain Ii.

Fig. 3B is a representation of a second minigene of the invention (SEQ ID NO: 148), in which the HLA-DRα chain leader peptide is linked to the amino terminus of a 24-residue blocking peptide fragment of human invariant 10 chain Ii.

### Experimental Data

#### METHODS

# I. Purification of HLA-DR antigens.

HLA-DR molecules were purified from homozygous, 15 Epstein-Barr virus-transformed, human B lymphoblastoid lines: DR1 from LG-2 cells, DR2 from MST cells, DR3 from WT20 cells, DR4 from Priess cells, DR7 from Mann cells, and DR8 from 23.1 cells. All of these cell lines are publicly available. Cell growth, harvest conditions and 20 protein purification were as previously described (Gorga, J. et al., 1991). Briefly, 200 grams of each cell type was resuspended in 10mM Tris-HCl, 1mM dithiothreitol (DTT), 0.1mM phenylmethylsulfonylflouride (PMSF), pH 8.0, and lysed in a Thomas homogenizer. The nuclei were 25 removed by centrifugation at 4000xg for 5 min and the pellets washed and repelleted until the supernatants were clear. All the supernatants were pooled and the membrane fraction harvested by centrifugation at 175,000xg for 40 The pellets were then resuspended in 10 mM Tris-30 HCl, 1mM DTT, 1mM PMSF, 4% NP-40. The unsolubilized membrane material was removed by centrifugation at 175,000xg for 2 hours, and the NP-40 soluble supernatant fraction used in immunoaffinity purification.

Detergent soluble HLA-DR was bound to a LB3.1protein A sepharose column (Gorga et al., <u>id</u>) and eluted
with 100 mM glycine, pH 11.5. Following elution, the
sample was immediately neutralized by the addition of
Tris-HCl and then dialyzed against 10mM Tris-HCl, 0.1%
deoxycholic acid (DOC). The LB3.1 monoclonal antibody
recognizes a conformational determinant present on the
nonpolymorphic HLA-DRa chain, and thus recognizes all
allotypes of HLA-DR.

The transmembrane domain of the DR molecules was removed by papain digestion, and the resulting water-soluble molecule further purified by gel filtration chromatography on an S-200 column equilibrated in 10mM Tris-HCl, pH 8.0. The purified DR samples were concentrated by ultrafiltration, yield determined by BCA assay, and analyzed by SDS polyacrylamide gel electrophoresis.

# II. Extraction and fractionation of bound peptides.

Water-soluble, immunoaffinity-purified class II 20 molecules were further purified by high-performance size exclusion chromatography (SEC), in 25 mM N-morpholino ethane sulfonic acid (MES) pH 6.5 and a flowrate of 1 ml/min., to remove any residual small molecular weight contaminants. Next, Centricon microconcentrators 25 (molecular weight cutoff 10,000 daltons) (Amicon Corp.) were sequentially washed using SEC buffer and 10% acetic acid prior to spin-concentration of the protein sample (final volume between 100-200  $\mu$ l). Peptide pools were extracted from chosen class II alleles by the addition of 30 1 ml of 10% acetic acid for 15 minutes at 70°C. These conditions are sufficient to free bound peptide from class II molecules, yet mild enough to avoid peptide degradation. The peptide pool was separated from the class II molecule after centrifugation through the

Centricon concentrator, with the flow-through containing the previously bound peptides.

The collected acid-extracted peptide pool was concentrated in a Savant Speed-Vac to a volume of 50  $\mu$ l 5 prior to HPLC separation. Peptides were separated on a microbore C-18 reversed-phase chromatography (RPC) column (Vydac) utilizing the following non-linear gradient protocol at a constant flowrate of 0.15 ml/min.: 0-63 min. 5%-33% buffer B; 63-95 min. 33%-60% buffer B; 95-105 10 min 60%-80% buffer B, where buffer A was 0.06% trifluoroacetic acid/water and buffer B was 0.055% trifluoroacetic acid/acetonitrile. Chromatographic analysis was monitored at multiple UV wavelengths (210, 254, 277, and 292 nm) simultaneously, permitting 15 spectrophotometric evaluation prior to mass and sequence analyses. Shown in Fig.1 are chromatograms for each of the six DR peptide pools analyzed. Collected fractions were subsequently analyzed by mass spectrometry and Edman sequencing.

# 20 III. Analysis of peptides.

The spectrophotometric evaluation of the peptides during RPC provides valuable information regarding amino acid composition (contribution of aromatic amino acids) and is used as a screening method for subsequent characterization. Appropriate fractions collected during the RPC separation were next analyzed using a Finnegan-MAT LaserMat matrix-assisted laser-desorption mass spectrometer (MALD-MS) to determine the individual mass values for the predominant peptides. Between 1%-4% of the collected fraction was mixed with matrix (1μ1 α-Cyano-4-hydroxycinnamic acid) to achieve mass determination of extracted peptides. The result of this analysis for HLA-DR1 is shown in Fig. 2. Next, chosen peptide samples were sequenced by automated Edman

degradation microsequencing using an ABI 477A protein sequencer (Applied Biosystems) with carboxy-terminal verification provided by mass spectral analysis using the Finnigan-MAT TSQ 700 triple quadruple mass spectrometer equipped with an electro-spray ion source. This parallel analysis ensures complete identity of peptide composition and sequence. Peptide alignment with protein sequences stored in the SWISS-PROT database was performed using the FASTA computer database search program. Set forth in Tables 1-10 are the results of this sequence analysis for each of the DR molecules studied.

## RESULTS

#### I. HLA-DR1.

The HLA-DR1 used in this study was papain 15 solubilized to enable the material to be used both for crystallographic and bound peptide analyses. The peptides bound to DR1 were acid extracted and fractionated using RPC (Fig. 1). The absence of any detectable peptidic material following a second 20 extraction/RPC separation verified quantitative peptide extraction. Amino acid analysis (ABI 420A/130A derivatizer/HPLC) of extracted peptide pools demonstrated a 70-80% yield, assuming total occupancy of purified DR1 with a molar equivalent of bound peptides corresponding 25 to the size distribution determined by mass spectrometry (see Fig. 2). The RPC profiles obtained from DR1 extractions of multiple independent preparations were reproducible. Furthermore, profiles from either detergent-soluble or papain-solubilized DR1 were 30 equivalent. To confirm that the peptides were in fact identical in detergent-soluble and papain-digested DR1, mass spectrometry and Edman sequencing analyses were performed and revealed identical masses and sequences for analogous fractions from the two preparations.

Matrix-assisted laser desorption mass spectrometry (MALD-MS) was used to identify 111 species of unique mass contained within the eluted peptide pool of DR1 with an average size of 18 and a mode of 15 residues (Fig. 2). 5 Over 500 additional mass species present within the molecular weight range of 13-25 residues were detected; however, the signal was not sufficient to assign individual masses with confidence. Multiple species of varying mass were detected in fractions corresponding to single RPC 10 peaks indicating co-elution of peptides. To characterize these peptides further, samples were analyzed in parallel on a triple quadruple mass spectrometer equipped with an electrospray ion source (ESI-MS) and by automated Edman degradation microsequencing (Lane et al., J. Prot. Chem. 15 10:151-160 (1991)). Combining these two techniques permits crucial verification of both the N- and C-terminal amino acids of peptides contained in single fractions. sequence and mass data acquired for twenty peptides isolated from DR1 are listed in Table 1. all the 20 identified peptides aligned with complete identity to regions of proteins stored in the SWISS-PROT database.

Surprisingly, sixteen of the twenty sequenced DR1-bound peptides were 100% identical to regions of the self proteins HLA-A2 and class II-associated invariant chain (Ii), representing at least 26% of the total extracted peptide mass. These isolated peptides varied in length and were truncated at both the N- and C-termini, suggesting that: 1) antigen processing occurs from both ends after binding to DR1, or 2) class II molecules bind antigen from a pool of randomly generated peptides. The yields from the peptide microsequencing indicated that HLA-A2 (Fig. 1) and Ii each represents at least 13% of the total DR1-bound peptides.

An additional surprising finding concerned a peptide 35 which, although bound to HLA-DR and 100% homologous with

HLA-A2 peptide, was derived from a cell which does not express HLA-A2 protein. Evidently this peptide is derived from a protein containing a region homologous with a region of HLA-A2 protein. Thus, for purposes of this specification, the term "HLA-A2 protein" is intended to include HLA-A2 protein itself, as well as any naturally occurring protein which contains a ten or greater amino acid long region of >80% homology with an HLA-DR-binding peptide derived from HLA-A2. An "HLA-A2 peptide" similarly refers to peptides from any HLA-A2 protein, as broadly defined herein.

The other four peptides identified in the DR1 studies were derived from two self proteins, transferrin receptor and the Na+/K+ ATPase, and one exogenous protein, 15 bovine serum fetuin (a protein present in the serum used to fortify the medium which bathes the cells). Each of these only 0.3-0.6% of the total peptides occupied population, significantly less than either the HLA-A2 or the Ii peptides. It is known that class II molecules en 20 route to the cell surface intersect the pathway of incoming endocytic vesicles. Both recycling membrane proteins and endocytosed exogenous protein travel this common pathway. Hence, the HLA-A2, transferrin receptor, Na+/K+ ATPase and bovine fetuin derived peptides would all encounter DR1 in Ii associates with nascent class II 25 a similar manner. molecules in the endoplasmic reticulum (ER) (Jones et al., Mol. Immunol. 16:51-60 (1978)), preventing antigen binding until the class II/Ii complex arrives at an endocytic (Roche and Cresswell, Nature 345:615-618 compartment 30 (1990)), where Ii undergoes proteolysis (Thomas et al., J. Immunol. 140:2670-2675 (1988); Roche and Cresswell, Proc. Natl. Acad. Sci. USA 88:3150-3154 (1991)), thus allowing peptide binding to proceed. Presumably, the Ii peptides bound to DR1 were generated at this step.

Synthetic peptides corresponding to five of the peptides reported in Table 1 were made and their relative binding affinities to DR1 determined. The influenza A hemagglutinin peptide (HA) 307-319 (SEQ ID NO: 24) has been 5 previously described as a high affinity, HLA-DR1 restricted peptide (Roche and Cresswell, J. Immunol. 144:1849-1856 (1990); Rothbard et al., Cell 52:515-523 (1988)), and was thus chosen as the control peptide. "Empty" DR1 purified from insect cells expressing recombinant DR1 cDNA was used 10 in the binding experiments because of its higher binding capacity and 10-fold faster association kinetics than DR1 isolated from human cells (Stern and Wiley, Cell 68:465-477 (1992)). All the synthetic peptides were found to compete well (Ki < 100 nM) against the HA peptide (Table 2). At 15 first approximation, the Ii 106-119 peptide (SEQ ID NO: 156) had the highest affinity of all the competitor peptides measured, equivalent to that determined for the control HA peptide. In addition to the Ki determinations, these peptides were found to confer resistance to 20 SDS-induced  $\alpha-\beta$  chain dissociation of "empty" DR1 when analyzed by SDS-PAGE, indicative of stable peptide binding (Sadegh-Nasseri and Germain, Nature 353:167-170 (1991); Dornmair et al., Cold Spring Harbor Symp. Quant. Biol. 54:409-415 (1989); Springer et al., J. Neither of the two control 25 252:6201-6207 (1977)). peptides,  $\beta_2$ m 52-64 (SEQ ID NO: 26) nor Ii 96-110 (SEQ ID 25), was able to either confer resistance to SDS-induced chain dissociation of DR1 or compete with HA 307-319 (SEQ ID NO: 24) for binding to DR1; both of these 30 peptides lack the putative binding motif reported in this study (see below).

A putative DR1 binding motif based on the sequence alignments of the core epitopes (the minimum length) of certain naturally processed peptides is shown in Table 3.

The peptides listed in this table include those determined

herein for HLA-DR1, as well as a number of peptides identified by others and known to bind DR1 (reference #6 in this table being O'Sullivan et al., J. Immunol. 145:1799-1808, 1990; reference #17, Roche & Cresswell, J. Immunol. 5 144:1849-1856, 1990; reference #25, Guttinger et al., Intern. Immunol. 3:899-906, 1991; reference #27, Guttinger et al. EMBO J. 7:2555-2558, 1988; and reference #28, Harris et al., J. Immunol. 148:2169-2174, 1992). The key residues proposed in the motif are as follows: a positively charged 10 group is located at the first position, referred to here as the index position for orientation (I); a hydrogen bond donor is located at I+5; and a hydrophobic residue is at I+9. In addition, a hydrophobic residue is often found at Every naturally processed peptide I+1 and/or I-1. 15 sequenced from DR1 conforms to this motif (with the exception of the HLA-A2 peptide 103-116 (SEQ ID NO: 3) that Because the putative motif is not lacks residue I+9). placed in a defined position with respect to the first amino acid and because of the irregular length of bound 20 peptides, it is impossible to deduce a motif from sequencing of peptide pools, as was done for class I molecules (Falk et al., Nature 351:290-296 (1991)). The Ii 96-110 peptide (SEQ ID NO: 25), a negative control peptide used in binding experiments, has the I and I+5 motif 25 residues within its sequence, but is missing eight additional amino acids found in Ii 105-118 (SEQ ID NO: 16) (Table 3C).

A sequence comparison of 35 previously described DR1-binding synthetic peptides (O'Sullivan et al., J. 30 Immunol. 145:1799-1808 (1990); Guttinger et al., Intern. Immunol. 3:899-906 (1991); Hill et al., J. Immunol. 147:189-197 (1991); Guttinger et al., EMBO J. 7:2555-2558 (1988); Harris et al., J. Immunol. 148:2169-2174 (1992)) also supports this motif. Of the 35 synthetic peptides, 21 (60%) have the precise motif, nine (30%) contain a single

shift at either I or I+9, and the remaining five (10%) have a single substitution at I (Table 3B and C). Interestingly, in the latter peptides, a positive charge at I is always replaced by a large hydrophobic residue (Table 8C); a 5 pocket has been described in class I molecules that can accommodate this precise substitution (Latron et al., Proc. Natl. Acad. Sci. USA 88:11325-11329 (1991)). Contributions by the other eight amino acids within the motif or the length of the peptide have not been fully evaluated and may 10 compensate for shifted/missing residues in those peptides Evaluation of the remaining 117 exhibiting binding. non-DR1 binding peptides cited in those studies (which peptides are not included in Table 3) indicates that 99 (85%) of these peptides do not contain the DR1 motif 15 proposed herein. Of the remaining 18 peptides (15%) that do not bind to DR1 but which do contain the motif, 6 (5%) are known to bind to other DR allotypes; the remaining 12 peptides may have unfavorable interactions at other positions which interfere with binding.

In contrast to the precise N-terminal cleavages 20 observed in the previous study of six peptides bound to the mouse class II antigen termed I-Ab and five bound to mouse I-Eb (Rudensky et al., Nature 3563:622-627 (1991)), the peptides bound to DR1 are heterogeneous at both the N- and In contrast to peptides bound to class I molecules, which are predominantly nonamers (Van Bleek and Nathenson, Nature 348:213-216 (1990); Rotzschke et al., (1990); Jardetzky et al., Nature Nature 348:252-254 353:326-329 (1991); Hunt et al., Science 255:1261-1263 30 (1992)), class II peptides are larger and display a high degree of heterogeneity both in length and the site of terminal truncation, implying that the mechanisms of processing for class I and class II peptides substantially different. Furthermore, the present results 35 suggest that class II processing is a stochastic event and

that a DR allotype may bind peptides of different lengths from a complex random mixture. The heterogeneity observed may be solely due to protection of bound peptides from further degradation. Thus, class II molecules would play 5 an active role in antigen processing (as previously proposed (Donermeyer and Allen, J. Immunol. 142:1063-1068 (1989)) by protecting the bound peptides from complete Alternatively, the predominance of 15mers degradation. bound to DR1 (as detected by both the MALD-MS and the 10 yields of sequenced peptides) could be the result of trimming of bound peptides. In any event, the absence of detectable amounts of peptides shorter than 13 and longer than 25 residues suggests that there are length constraints intrinsic either to the mechanism of peptide binding or to 15 antigen processing. The predominance of peptides bound to from endogenously synthesized that are derived proteins, and particularly MHC-related proteins, may result from the evolution of a mechanism for presentation of self peptides in connection with the generation of self 20 tolerance.

# II. Other HLA-DR molecules.

The sequences of naturally processed peptides eluted from each of DR2, DR3, DR4, DR7 and DR8 are shown in Tables 4-8, respectively. In addition to those peptides shown in Table 4, it has been found that DR2 binds to long fragments of HLA-DR2a β-chain and HLA-DR2b β-chain, corresponding to residues 1-126 or 127 of each of those proteins. Presumably, only a short segment of those long fragments is actually bound within the groove of DR2, with the remainder of each fragment protruding from one or both ends of the groove. Table 9 gives sequences of DR1 from another cell line which does not have wild-type Ar, but which has bound A2-like peptides. Table 10 gives sequences of peptides eluted from DR4 and DR11 molecules expressed in

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cells from a human spleen. These data demonstrate the great prevalence of self peptides bound, compared to exogenous peptides. The data also show that the A2 and Ii peptides occur repeatedly. In addition, certain of the Tables include peptides that appear to derive from viral proteins, such as Epstein-Barr virus major capsid protein, which are likely to be present in the cells studied.

# III. Peptide Delivery Genetic Constructions.

In order to prepare genetic constructs for <u>in vivo</u> administration of genes encoding immunomodulatory peptides of the invention, the following procedure is carried out.

Overlapping synthetic oligonucleotides were used to generate the leader peptide/blocking peptide mini-genes illustrated in Fig. 3 by PCR amplification from human HLA-DRα and invariant chain cDNA templates. These mini-genes encode the Ii peptide fragments KMRMATPLLMQALPM (or Ii<sub>15</sub>; SEQ ID NO: 15) and LPKPPKPVSKMRMATPLLMQALPM (or Ii<sub>24</sub>; SEQ ID NO: 7). The resulting constructs were cloned into pGEM-2 (Promega Corp.) to form the plasmids pGEM-2-α-Ii<sub>15</sub> and pGEM-2-α-Ii<sub>24</sub>, with an upstream T7 promoter for use in the <u>in vitro</u> transcription/translation system described below.

in vivo expression, each mini-gene was subsequently subcloned from the pGEM-2 derivatives into a 25 transfection vector, pHβactin-1-neo (Gunning et al., (1987) Proc. Natl. Acad. Sci. U.S.A. 84:4831), to form the plasmids  $pH\beta$ actin- $\alpha$ - $Ii_{15}$  and  $pH\beta$ actin- $\alpha$ - $Ii_{24}$ . The inserted thus expressed <u>in vivo</u> from mini-genes are constitutive/strong human  $\beta$  actin promoter. In addition, 30 the mini-genes were subcloned from the pGEM-2 derivatives into the vaccinia virus recombination vector pSC11 (S. Chakrabarti et al. (1985) Mol. Cell. Biol. 5, 3403-3409) to form the plasmids pSC11- $\alpha$ -Ii<sub>15</sub> and pSC11- $\alpha$ -Ii<sub>24</sub>. Following recombination into the viral genome the inserted mini-genes are expressed from the strong vaccinia  $p_{7.5}$  promoter.

# Intracellular trafficking signals added to peptides.

Short amino acid sequences can act as signals to 5 target proteins to specific intracellular compartments. For example, hydrophobic signal peptides are found at the amino terminus of proteins destined for the ER, while the sequence KFERQ (SEQ ID NO: 153) (and other closely related sequences) is known to target intracellular polypeptides to 10 lysosomes, while other sequences target polypeptides to endosomes. In addition, the peptide sequence KDEL (SEQ ID NO: 152) has been shown to act as a retention signal for the ER. Each of these signal peptides, or a combination thereof, can be used to traffic the immunomodulating 15 peptides of the invention as desired. For example, a construct encoding a given immunomodulating peptide linked to an ER-targeting signal peptide would direct the peptide to the ER, where it would bind to the class II molecule as it is assembled, preventing the binding of intact Ii which 20 is essential for trafficking. Alternatively, a construct can be made in which an ER retention signal on the peptide would help prevent the class II molecule from ever leaving If instead a peptide of the invention is targeted to the endosomic compartment, this would ensure that large 25 quantities of the peptide are present when invariant chain is replaced by processed peptides, thereby increasing the likelihood that the peptide incorporated into the class II complex is the high-affinity peptides of the invention rather than naturally-occurring, potentially immunogenic The likelihood of peptides of the invention 30 peptides. being available incorporation into class II can increased by linking the peptides to an intact Ii Since Ii is known to traffic polypeptide sequence. class II molecules to the endosomes, the hybrid Ii would

carry one or more copies of the peptide of the invention along with the class II molecule; once in the endosome, the hybrid Ii would be degraded by normal endosomal processes to yield both multiple copies of the peptide of the invention or molecules similar to it, and an open class II binding cleft. DNAs encoding immunomodulatory peptides containing targeting signals will be generated by PCR or other standard genetic engineering or synthetic techniques, and the ability of these peptides to associate with DR molecules will be analyzed in vitro and in vivo, as described below.

It is proposed that the invariant chain prevents class II molecules from binding peptides in the ER and may contribute to heterodimer formation. Any mechanism that prevents this association would increase the effectiveness of class II blockade. Therefore, a peptide corresponding to the site on Ii which binds to the class II heterodimer, or corresponding to the site on either the  $\alpha$  or  $\beta$  subunit of the heterodimer which binds to Ii, could be used to prevent this association and thereby disrupt MHC class II function.

# In Vitro Assembly.

cell free extracts are used routinely for expressing eukaryotic proteins (Krieg, P. & Melton, D. (1984) Nucl. Acids Res. 12, 7057; Pelham, H. and Jackson, R. (1976) Eur. J. Biochem. 67, 247). Specific mRNAs are transcribed from DNA vectors containing viral RNA polymerase promoters (Melton, D. et al. (1984) Nucl. Acids Res. 12, 7035), and added to micrococcal nuclease-treated cell extracts. The addition of 35s methionine and amino acids initiates translation of the exogenous mRNA, resulting in labeled protein. Proteins may be subsequently analyzed by SDS-PAGE and detected by autoradiography. Processing events such as signal peptide cleavage and core glycosylation are

initiated by the addition of microsomal vesicles during translation (Walter, P. and Blobel, G. (1983), Meth. Enzymol., 96, 50), and these events are monitored by the altered mobility of the proteins in SDS-PAGE gels.

The ability of peptides containing a signal peptide sequence to be accurately processed and to compete with invariant chain for class II binding in the ER are assayed in the in vitro system described above. Specifically, DR1  $\alpha$ - and  $\beta$ -chain and invariant chain peptide constructs 10 described above are transcribed into mRNAs, which will be translated in the presence of mammalian microsomal membranes. Association of the DR heterodimer with Ii is determined by immunoprecipitation with antisera to DR and Ii. Addition of mRNA encoding the peptide of the invention 15 to the translation reaction should result in a decreased level of coimmunoprecipitated Ii, and the concomitant appearance of coimmunoprecipitated peptide, as determined by SDS-PAGE on TRIS-Tricine gels. These experiments will provide a rapid assay system for determining the potential 20 usefulness of a given blocking peptide as a competitor for Ii chain binding in the ER. Those peptides of the invention which prove to be capable of competing successfully with Ii in this cell-free assay can then be tested in intact cells, as described below.

# 25 In Vivo Assembly.

Human EBV-transformed B cell lines LG-2 and HOM-2 (homozygous for HLA-DR1) and the mouse B cell hybridoma LK35.2 are transfected with either  $50\mu g$  of linearized pH $\beta$ actin- $\alpha$ -Ii $_{15}$  or pH $\beta$ actin- $\alpha$ -Ii $_{24}$  or (as a control) 30 pH $\beta$ actin-1-neo by electroporation (150mV, 960 $\mu$ F, 0.2cm cuvette gap). Following electroporation, the cells are cultured in G418-free medium until total recovery (approximately 4 days). Each population is then placed under G418 selection until neomycin-expressing resistant

populations of transfectants are obtained (approximately 1-2 months). The resistant populations are subcloned by limiting dilution and the clonality of stable transfectants determined by PCR amplification of blocking peptide mRNA expression.

Stable transfectants of LG-2 and HOM-2 carrying blocking peptide mini-genes or negative control vectors are grown in large-scale culture conditions until 20 grams of pelleted cell mass is obtained. The HLA-DR expressed by each transfectant is purified, and the bound peptide repertoire (both from within the cell and from the cell surface) analyzed as described above. Successful demonstration of a reduction in the total bound peptide diversity will be conclusive evidence of intracellular delivery of immuno-modulatory peptides.

utilizes cell-based assay second transfectants of LK35.2 cells carrying blocking peptide mini-genes or negative control vectors; these cells are used as APCs in T cell proliferation assays. 20 transfectant is cultured for 24 hours in the presence of different dilutions of hen egg lysozyme (HEL) and HELspecific T cell hybridomas. The relative activation of the T cells present in each assay (as measured by lymphokine production) is determined using the publicly available 25 lymphokine dependent cell line CTLL2 in a 3H-thymidine incorporation assay (Vignali et al. (1992) J.E.M. 175:925-Successful demonstration of a reduction in the ability of blocking peptide expressing transfectants to present HEL to specific T cell hybridomas will 30 conclusive evidence of intracellular delivery of immunomodulatory peptides. Cells of the human TK cell line 143 (ATCC) are infected with vaccinia virus (strain WR, TK+) (ATCC), and two hours postinfection, pSC11- $\alpha$ -Ii<sub>15</sub> or pSC11- $\alpha\text{-Ii}_{24}$  or pSC11 is introduced into the infected cells by 35 calcium phosphate precipitation. TK recombinants are

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selected for with bromodeoxyuridine at  $25\mu g/ml$ . Recombinant plaques are screened by PCR for the presence of mini-gene DNA. Recombinant virus is cloned by three rounds of limiting dilution to generate pure clonal viral stocks.

In experiments analogous to the transfection experiments described above, recombinant vaccinia viruses encoding mini-genes or vector alone will be used to infect large-scale cultures of the human EBV transformed B cell lines LG-2 and HOM-2. Following infection, the HLA-DR is purified and the bound peptide repertoire analyzed as described above. A reduction of the complexity of the bound peptide population and a significant increase in the relative amount of Ii peptides bound are conclusive evidence that vaccinia can deliver blocking peptides to human APCs.

The same recombinant vaccinia viruses encoding minigenes or vector will be used to infect mice experiencing experimentally-induced autoimmunity. A number of such models are known and are referred in Kronenberg, Cell 65:537-542 (1991).

# Liposomal Delivery of Synthetic Peptides or Mini-gene Constructs.

Liposomes have been successfully used as drug carriers and more recently in safe and potent adjuvant strategies for malaria vaccination in humans (Fries et al. (1992), Proc. Natl. Acad. Sci. USA 89:358). Encapsulated liposomes have been shown to incorporate soluble proteins and deliver these antigens to cells for both in vitro and in vivo CD8<sup>+</sup> mediated CTL response (Reddy et al., J. Immunol. 148:1585-1589, 1992; and Collins et al., J. Immunol. 148:3336-3341, 1992). Thus, liposomes may be used as a vehicle for delivering synthetic peptides into APCs. Harding et al. (Cell (1991) 64, 393-401) have

demonstrated that the targeting of liposome-delivered

antigen to either of two intracellular class II-loading compartments, early endosomes and/or lysosomes, can be accomplished by varying the membrane composition of the liposome: acid-sensitive liposomes were found to target their contents to early endosomes, while acid-resistant liposomes were found to deliver their contents to lysosomes. Thus, the peptides of the invention will be incorporated into acid-sensitive liposomes where delivery to endosomes is desired, and into acid-resistant liposomes for delivery to lysosomes.

Liposomes are prepared by standard detergent dialysis or dehydration-rehydration methods. dioleoylphosphatidylethanolamine liposomes, (DOPE) and palmitoylhomocystein (PHC) are utilized, while 15 dioleoylphospatidylcholine (DOPC) dioleoylphosphatidylserine (DOPS) are used for preparation of acid-resistant liposomes. 10<sup>-5</sup> mol of total lipid (DOPC/DOPS or DOPE/PHC at 4:1 mol ratios) are dried, hydrated in 0.2 ml of HEPES buffered saline (HBS) (150 mM 20 NaCl, 1 mM EGTA, 10mM HEPES pH 7.4) and sonicated. lipid suspensions are solubilized by the addition of 0.1 ml of 1 M octylglucoside in HBS. The peptides to be entrapped are added to 0.2 ml of 0.6 mM peptide in 20% HBS. mixture is then frozen, lyophilized overnight, liposomes will be treated with These 25 rehydrated. chymotrypsin to digest any surface-bound peptide. Liposome delivery to EBV-transformed cell lines (as described above) will be accomplished by 12-16 hour incubation at 37°C. HLA-DR will be purified from the liposome treated cells and 30 bound peptide analyzed as above.

Alternatively, the liposomes are formulated with the DNA mini-gene constructs of the invention, and used to deliver the constructs into APCs either <u>in vitro</u> or <u>in vivo</u>.

Human immunization will be carried out under the protocol approved by both The Johns Hopkins University Joint Committee for Clinical Investigation and the Human Subject Research Review Board of the Office of the Surgeon 5 General of the U.S. Army (Fries et al. (1992), Proc. Natl. Acad. Sci. U.S.A. 89:358-362), using dosages described therein, or other dosages described in the literature for liposome-based delivery of therapeutic agents.

## Delivery via Immune-stimulating Complexes (ISCOMS).

10 ISCOMS are negatively charged cage-like structures 30-40nm in size formed spontaneously on mixing cholesterol and Quil A (saponin). Protective immunity has been generated in a variety of experimental models of infection, including toxoplasmosis and Epstein-Barr virus-15 induced tumors, using ISCOMS as the delivery vehicle for antigens (Mowat and Donachie) Immunology Today 12:383-385, Doses of antigen as low as  $1\mu g$  encapsulated in ISCOMS have been found to produce class I mediated CTL responses, where either purified intact HIV-1-IIIB gp 160 20 envelope glycoprotein or influenza hemagglutinin is the antigen (Takahashi et al., Nature 344:873-875, 1990). Peptides are delivered into tissue culture cells using ISCOMS in a manner and dosage similar to that described above for liposomes; the class II peptide binding of 25 delivered peptides are then determined by extraction and characterization as described above. ISCOM-delivered peptides of the invention which are effectively utilized by cultured cells are then tested in animals or humans.

In addition to delivery of the therapeutic synthetic peptides, ISCOMS could be constituted to deliver the minigene constructs to APCs, and thus serve as an alternative to the above-outlined vaccinia strategy.

# Immunogenic Peptide Delivery (Vaccines).

using the above-described In addition to intracellular delivery systems to deliver nonimmunogenic self peptides with the specific aim of down-modulating the 5 immune system (thus alleviating autoimmune conditions), the delivery systems of the invention may alternatively be used as a novel means of vaccination, in order to stimulate a portion of the immune system of an animal. In the latter context, the delivery system is employed to deliver, into 10 appropriate cells, DNA constructs which. express peptides pathogen-derived intended immunogenic, stimulate an immune response against a specific pathogen. Because the antigenic peptide is produced inside the target cell itself, the vaccine method of the invention ensures 15 that there is no circulating free antigen available to stimulate antibody formation and thereby induce potentially deleterious or inappropriate immunological reactions. immune response stimulated by vaccines of the invention is, because the vaccines are targeted solely to APC's, limited 20 to the T cell mediated response, in contrast to standard vaccine protocols which result in a more generalized immune response. Although some of the peptide-presenting APC's will initially be lysed by host T cells, such lysis will be limited because, inter alia, the virus-based vaccine is 25 non-replicative, i.e., each carrier virus can infect only one cell.

The model antigen that will be used to perfect and test the system of the invention is hen egg lysozyme (HEL). It is arguably the most well characterized protein for antigen presentation studies, to which there are numerous monoclonal antibodies and class I- and class II-restricted mouse T cell clones and hybridomas. The primary epitopes that will be studied are the peptide HEL 34-45, as both monoclonal antibodies and CD4+ T cell hybridomas are available, and peptide HEL 46-61, as both class I and class

II-restricted T cell clones and hybridomas have been raised and are publicly available. These two sequences are thus proven immunogenic epitopes. Initially, four constructs encoding different polypeptides are analyzed: (a) whole, 5 secreted HEL, (B) HEL 34-45, (c) HEL 46-61, and (d) HEL 34-The last three include a signal sequence known to be cleaved in these cells, e.g., IAk (MPRSRALILGVLALTTMLSLCGG; ), which would result in targeting to the ER. All constructs are then subcloned into pH\$Apr-1 neo. 10 methodology for making these constructs is similar to that The constructs are introduced into outlined above. appropriate APCs, e.g., LK35.2 cells, by means of a conventional eukaryotic transfection or one of the delivery vehicles discussed above (e.g., vaccinia, liposomes, or 15 ISCOMS). LK35.2 cells, which possess the mouse MHC Class II restriction molecules IAk and IEk, transfected with each of the constructs are tested for their ability to stimulate the appropriate class I and class II-restricted T cell hybridomas and clones using standard techniques. Whether 20 class I stimulation is observed will depend on whether peptide trimming can occur in the ER, in order to produce an 8-10-mer suitable for binding to class I molecules. If these constructs are ineffective for class I stimulation, they can be modified in order to produce a more effective 25 peptide for class I binding. If these constructs prove to be less effective for class II-restricted responses, they can be tagged with endosomal and/or lysosomal targeting sequences as discussed in Section V.

The effectiveness of targeting signals used to direct immunogenic peptides to particular intracellular organelles would be monitored using electron microscopic analysis of immunogold stained sections of the various transfectants. Rabbit anti-peptide antisera would be produced and affinity purified for this application. In

addition, monoclonal antibody HF10, which recognizes HEL 34-45, will be used.

Once a construct is defined that can be effectively presented by transfectants in vitro, its effectiveness in vivo will be determined. This can be tested by injection of the transfectants i.p. and/or s.c. into C3H/Balb/c Fl mice, or by injection of the construct incorporated into an appropriate delivery vehicle (e.g., liposome, ISCOMS, retrovirus, vaccinia). Optimal protocols and doses for such immunizing injections can be determined by one of ordinary skill in the art, given the disclosures provided herein. Efficiency of immunization can be tested by standard methods such as (a) proliferation of class II-restricted T cells in response to HEL pulsed APCs, (b) CTL response to 51Cr-labeled targets, and (c) serum antibody titre as determined by ELISA.

Once the details of the vaccine delivery system of the invention are optimized, constructs encoding peptides with useful immunizing potential can be incorporated into Such peptides can be identified by standard 20 the system. means now used to identify immunogenic epitopes on pathogen-derived proteins. For example, candidate peptides for immunization may be determined from antibody and T cell analysis of animals infected with a particular pathogen. 25 In order to obtain a protective and effective anamnestic response, the peptides used for vaccination should ideally be those which are presented with the highest frequency and efficiency upon infection. This could best be determined by using the procedures outlined in the experimental 30 section above to extract and characterize the peptides bound by MHC class II molecules from infected cells. Given allelic restriction of immunogenic peptides (in contrast to the observed degenerate binding of self peptides of invention), a mini-gene encoding several immunogenic 35 peptides will probably be required to provide a vaccine WO 94/04171 PCT/US93/07545

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useful for the entire population. Vaccine administration and dosage are as currently employed to smallpox vaccination.

TABLE 1 LG-2/HLA-DRT BINDING PEPTIDES

PROTEIN SOURCE	P051110H	SEGLENCE	SEQ 10 110.	LENGTH	FRACTION	3	MASS SPEC	71EG	
Paeudo HLA-A2	103-120	VGSDURFLRGYHQYAYDG	-	18	DR15-59	2190.4	2190.4	39.5	l
	103-117	VGSDWRFLRGYHQYA	7	<b>\$</b>	DR15-58	1855.0	1854.4	\$.709	
	103-116	VGSDLRFLRGYHOY	m	75	DR15-58	1784.0	1783.6	53.3	
	104-117	GSDURFLRGYHQYA	4	14	DR15-56	175.3	175.2	96.5	
	105-117	SDURFLEGYHOYA	'n	13	DR15-56	1698.2	1698.8	8.87	
	97-121	I PKPPKPVSKMRMATPLLMGALPMG	•	×	DR15-88	2733.5	2734.5	\$.05	
INVALIBILITY CHESTIN	97-120	LPKPPKPVSKHRMATPLLHDALPH	1	72	DR15-88	2676.4	2675.9	80.8	
	98-121	PKPPKPVSKHRMATPLLMQALPMG	80	%	DR15-86	2,0292	2619.7	91.5	
	97-119	LPKPPKPVSKMRMATPLLMGALP	٥	23	<b>0R1S-86</b>	2.245.2	2544.5	112.2	
	98-120	PKPPKPVSKHRNATPLLMGALPM	5	8	DR15-87	2563.2	2562.3	145.0	•
	99-120	KPPKPVSKHRMATPLLHOALPH	=	23	DR15-87	2466.1	2465.8	101.5	3 (
	98-119	PKPPKPVSKMRMATPLLMGALP	12	22	DR15-84	2432.0	2431.7	72.5	5
	99-119	KPPKPVSKMRMATPLLMGALP	<b>.</b>	72	DR15-84	2334.9	2334.2	31.6	
	100-119	PPKPVSKMRMATPLLMGALP	71	20	<b>0R1S-86</b>	2506-7	2207.4	89.8	
	106-120	KKRMATPLLMGALPM	<b>\$</b>	15	DR15-86	1732.2	1731.9	178.5	
	106-119	KHRMATPLLMAALP	45	7	DR15-86	1601.0	1600.2	162.0	
Wa+/K+ ATPose	199-216	IPADLRIISANGCKYDNS	11	18	DR15-56	1686.6	1865.8	8.87	
Transferrin Recpt.	969-089	RVETHFLSPTVSPKESP	18	17	0815-58	2035.3	2036.8	30.3	
	72-95	YKHTLNGIDSVKWPRRPT	19	19	DR15-51	2237.6	2236.5	0,99	
	56-73	YKHTLNGIDSVKVAPRRP	02	81	DR15-50	2338.7	2338.5	32.5	
W. B. DR. A.chain	43-61	DVGEYRAVTELGRPDAEYU	21	4	<b>DR15-51</b>	228.5	~		
Carboxypept idase E	101-115	EPGEPEFKYICHMHG	22	<b>\$</b>	DR15-48	1704.9	1700.4		
				•			ESI-MS		

TABLE 2
PEPTIDE BINDING TO MLA-DR1

PEPTIDE	SEG 10 NO.	LENGTH	Ki vs IIA 307-319 <sup>b</sup>	SDS-Resistance <sup>c</sup> nM
			74	
HLA-A2 103-117	2	15	£ \$ 67	٠
11 105-119	\$1	15	o 10	•
11 97-120	^	72	33 1 5	•
Na+/K+ ATPase 199-216	11	18	6 1 89	•
Transf. Recept. 680-696	18	17	ot >	•
Bovine fetuin 56-72	25	19	81 18	•
HA 307-319	%	7	< 10	•
11 97-111	8	51	, 10 <b>,</b>	•
β <sub>2</sub> m 52-64	%	13	> 10 <b>*</b>	,

The first six entries correspond to peptides found associated with MLA-DR1 and the sequences are shown in Table 1. Two control peptides were also tested: \$2.64, SDLSFSKOWSYTL, is from human \$2.microglobulin and 11 96-110, LPKPPKPVSKMRMAI is a truncated version of the longest invariant chain derived peptide isolated from HLA-DR1. Peptides were synthesized using solid-phase Fmcc chemistry, deprotected and cleaved using standing methods, then purified by RPC. Purified peptides were analzyed by mass spectrometry and concentrations were determined by quantitative ninhydrin enelysis.

inhibition constants (Ki) were measured as the concentration of test peptide which inhibited 50% of the <sup>125</sup>1-labeled MA 307-319 binding to "empty" MLA-DRI produced in 519 insect cells (20). MA 307-319 was tabeled using Na<sup>12</sup>1 and chloramine-1 and isolated by gel filtration. Specific activity, determined by BCA assay (Pierce) and gamma counting, was 26,000 cpm/pmol. 10M labeled peptide and 10 mM purified MLA-DRI were mixed with 10 different concentration in the 10 pM) of synthetic cold competitor peptide in phosphate-buffered asline, pM 7.2, containing 1 mM E01A, and PMS of 10.1 mM lodoscetamide, and 3 mM MaM<sub>3</sub>, and incubated at 37°C for 85 hours. Free and bound peptide were separated by native gel electrophoresis (33) and bound radioactivity was quantitated using a Fujix inaging plate analyzer (8AS 2000) after four hour exposures on the phospho-imaging plates. Percent inhibition was calculated as the ratio of background-corrected radioactivity in a parallel sample containing no competitor peptide. Under these conditions, Ki measurements < 10 rM could not be accurately

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The ability of the synthetic peptides to confer resistence to SDS-induced chain dissociation of HLA-DRI produced in insect cells was determined as described (20). Briefly, 20 µH MLA-DRI was incubated with five-fold excess of synthetic peptide at 37°C for 85 hours, in phosphate-buffered saline (PH 7.2) with the protease inhibitor mixture described above. After incubation, the samples were analyzed by SDS-PAGE with and without boiling prior to loading. Peptides which prevented SDS-induced chain dissociation are indicated positive (\*) and those that did not negative

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TABLE 3 - PUTATIVE HLA-DR1 PEPTIDE BINDING MUTIF

A PROTEIN SOURCE	PEPTIDE SEGUENCE	SEQ 10 NO.	LENGTH		ACT ENEMOS.
HLA-A2	SOVRFLAGYHOYA	2	13	105-117	This study
Inverient Chain	KMRMAIPLLMAALP	16	14	105-118	
Ma+/K+ ATPase	I PADLR I I SANGCKVONS	11	18	199-216	
Transferrin Receptor	RVEYHFLSPTVSPKESP	18,	17	969-089	
Bovine Fetuin	YKHTLHQIOSYKVAPRRP	20	18	56-73	
13H 6	KVFGRCELAAAMKRHGLD	22	18	1-18	•
	RHRCKGTDVOAMIRGCRL	28	18	112-129	٠
E-	HPPHIEIOMLKNGKKI	62	16	31-46	ý
	NELGREKHTOACCRTH	30	16	19-34	9
7	SKPKVYQUFOLRKY	31	72	115-128	•
MASE	ATSTKKLHKEPATLIKAIDG	32	20	1-20	•
	PATLIKAIDGDTVKLMYKGO	33	50	11-30	•
	DRVKLMYKGOPMTFRLLLVD	34	20	21-40	9
Ϋ́Α	VAYVYKPHNTHEGHLRKSEA	35	20	111-130	9
HIV 013	OKOEPIDKELYPITSL	36	5	97-112	٠
HIV 017	GARASVLSGGELDKWE	37	51	1-16	9
Influenza KA	RTLYQNOGTYVSVGTSTLHK	38	20	187-206	•
Influenza HA	PKYVKQHTLKLAT	72	13	307-319	. 21
p (atcip. p190	LKKLVFGYRKPLONI	39	15	249-263	22
P falcip. CS	KHIEQYLKKIKNS	07	13	329-341	27
chicken OVA	DVFKELKVHHANENIE	13	16	15-30	•
081 A chain	CDTRPRFLWOLKFECHFFWG	27	20	1-20	28
	TERVRLLERCIYNOEESVRFOS	53	22	21-12	82
	DILEGRRAAVDTYCRHNYGVGESFT	3	22	∙ 06-99	28
2	KAERADLIAYLKGATAK	\$\$	11	88 - 104	•
and along the state of	CRICOENPOVHFFKNIVIPRIPPP	. 97	72	75-98	9

Table 3, continued

A PROTEIN SOURCE	PEPTIDE SEGUENCE	SEG 10 MG.	LENGTH	POSITION	REFERENCE
C Influenza Matrix	PLKAETAORLEDV	23	13	16-31	9
HIV 017	ROTLGOLOPSLOTGSE	85	91	57-72	•
	INDAMENTAL INTERPRETATION OF THE PROPERTY OF T	67	16,	7-22	ø
	INTRCYKLENPVIGCG	20	16	85-100	9
0 felcin p100	YKLMFYFDLLRAKL	51	71	211-224	52
and the second	IDTLKKNENIKEL	25	13	338-350	22
niet, & ton	DVCEYRAVIELGRPDAEYWN	23	20	79-65	28
urv 017	EBEAVAPOLLETSECC	×	91	41-56	•9
	DNYBCYSI GNUVCAAKFESHFTO	23	23	20-42	•
u	EALVROGLAKVATVYKPINT	95	20	101-120	•
25	PIVONLOGOMVHOAIS	57	5	1-16	•
2	SAI SEGATPODINIME	28	91	41-56	•
( <b>Q</b>	SFYILANTEFTPTETD	29	91	91-19	۰
25 a	KMYFWLIMIKCYKLEM	09	16	70-62	9

TABLE 4
MST/KLA-DR2 BINDING PEPTIES

Pseudo MLA-A2 103-120 103-119 104-119 105-119 105-119 105-117	03-120 03-119 04-119 03-117 03-116 105-117 97-120	VGSDVRFLRGYHOYAYDG VGSDVRFLRGYHOYAYD GSDVRFLRGYHOYA VGSDVRFLRGYHOYA VGSDVRFLRGYHOYA GSDKRFLRGYHOYA GSDKRFLRGYHOYA SOURFLRGYHOYA RPPREVSKYRNATPLLHOALPH RKPPKPVSKYRNATPLLHOALPH RKPPKPVSKYRNATPLLHOALPH RKPPKRVSKYRNATPLLHOALPH	1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	8 7 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	DR2-3-57 DR2-3-57 DR2-3-56 DR2-3-56 DR2-3-56 DR2-3-56	2190.4 2133.3 2034.3	2189.0 2131.8 2040.4 1858.5
<u> </u>	119 1119 1117 1117 120	VGSDIRFLRCTHOYATAD GSDIRFLRCTHOYATO VGSDIRFLRCTHOYA VGSDIRFLRCTHOYA GSDURFLRCTHOYA LPKPPKPVSKHRMATPLLHQALPH KPPKPVSKHRMATPLLHQALPH KPPKPVSKHRMATPLLHQALPH RPPKPVSKHRMATPLLHQALPH	2 2 2 5 2 5 2 1 2 2 5 2 1 2 2 2 2 2 2 2	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	DR2-3-57 DR2-3-56 DR2-3-56 DR2-3-56	2153.3 2034.3	2131.6 2040.4 1858.5
rient Chein	119 116 117 117 120	GSDURFLRCTHOTA VGSDURFLRCTHOTA VGSDURFLRCTHOTA SDURFLRCTHOTA SDURFLRCTHOTA LPKPPKPVSKORMATPLLHOALPH KPPKPVSKORMATPLLHOALPH KPPKPVSKORMATPLLHOALPH	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	\$ £ 7 7 E % E 8 8 8	DR2-3-56 DR2-3-56 DR2-3-56 DR2-3-55	2034.3	2040.4
rient Chein	117 117 117 120	VGSDIRFLRCYHOYA VGSDIRFLRCYHOYA GSDIRFLRCYHOYA SOLRFLRCYHOYA LPKPPKPVSKNRNATPLLHOALPH KPPKPVSKNRNATPLLHOALPH KPPKPVSKNRNATPLLHOALPH PKPPKRVSKNRNATPLLHOALPH	2 2 3 5 5 2 5	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	DR2-3-56 DR2-3-56 DR2-3-55		1858.5
rient Chein	116 117 117 120	VGSDURFLRCTHOY GSDURFLRCTHOYA SOURFLRCTHOYA LPKPPKPVSKHRMIPLLHOALPH KPPKPVSKHRMIPLLHOALPH KPPKPVSKHRMIPLLHOALPH KPPKPVSKHRMIPLLHOALPH PKPPKRVSKHRMIPLLHOALPH		2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	DR2-3-56 DR2-3-55	1855.0	
rient Chein	117 117 120 120	GSDURFLECTHOTA SOURFLECTHOTA LPKPPKPVSKHRMIPLLHOALPH RPPKPVSKHRMIPLLHOALPH KPPKPVSKHRMIPLLHOALPH RPPKRVSKHRMIPLLHOALPH	, , s , s , s , s , s , s , s , s , s ,	* 5 % 5 % 8 %	DR2-3-55	1784.0	1786.3
rient Chein	120	SOURFLECTHOYA LPKPPKPVSKHRMATPLLHGALPH PKPPKPVSKHRMATPLLHGALPH KPPKPVSKHRMATPLLHGALPH PKPPKPVSKHRMATPLLHGALPH		2 2 2 2 2	F C C C	1755.3	1755.0
rient Chein	120 120	LPKPPKPVSKYRNATPLLMAALPH PKPPKPVSKYRNATPLLMAALPH KPPKPVSKYRNATPLLMAALPH PKPPKPVSKYRNATPLLMALPH	, <b>,</b> 0 =	% % % %	DRZ-3-36	1698.2	1702.6
rient Chein	5 2	PKPPKPVSKARMATPLLMOALPH KPPKPVSKARMATPLLMOALPH PKPPKPVSKARMATPLLMOALP	. 2 =	2 2 2	DR2-3-70	2676.4	2675.0
	2	KPPKPVSKKRMATPLLMALPM KPPKPVSKRRMATPLLMALPM	: =	2 22	0R2-3-70	2563.2	2562.0
<b>*</b>	60,	PKPPKPVSKORNATPLLMGALP		22	0R2-3-70	2466.1	2465.0
•	02.	FRFFFF	: 2		082-3-66	2432.0	2437.0
-84	411-94 00-114	O SACRETOTAMENTOLOGICA	: =	12	DR2-3-66	2334.9	2340.0
	A11.44	AFTER SACRET LEGISTE	: 5	20	082-3-70	2206.7	2207.0
A11-001	<u> </u>	PERFESHINATION CONTRACTOR	: 3	61	0R2-3-71	2070.5	2074.3
371 - 901	<b>7</b>	SABATEL WALES	; <u>~</u>	: 52	082-3-70	1732.2	1732.0
	021.001	NAME OF TAXABLE OF TAX	158	ສ	DR2-3-44	2476.8	2478.1
HLA-DQ a-chein yr- 97-	97-112	MIVIKASHSTAATHEV	159	<b>1</b> 5	DR2-3-41	1716.9	1717.0
	;		140	81	DR2-3-41	1917.1	1920.5
HLA-DQ Ø-chain 42-	45-24		191	- 11	DR2-3-41	1830.0	1633.3
VC-64	<u>,</u>		162	. <b>2</b> 2	DR2-3-41	1629.8	1632.9
	, i		163	. £1	DR2-3-36	1353.5	1362.0
HLA-DR e-chein 162-	182-194	ANDELE CHAN	2 2	11	DF2-3-41	1697.9	1701.0
184-197 (MET) Kinase-relate 59-81 trasforming protein	59-81	ENNIFLGATNY I YVLMEEDLOKY	<b>\$</b>	8	DR2-3-65	2746.1	2746.6
	03/-/4/	OF! KWKYYOVPRKG! DA	8	11	DR2-3-71	2063.4	2074.3
Guarrylate-bind.	174-193	IQNL IKEEAFLGI TDEKTEG	19	50	DR2-3-70	5.848.5	2248.0*

Table 4, continued

PROTEIN SOURCE	POSITION	SEGLENCE	SEQ 10 NO.	LENCTH	FRACTION	₹	MSS SPEC
	1200-1220	SDKKI HIYAMII I DROVPOTO	\$91	21	DR2-3-61	2484.8	5490.9
Apol ipoprotein 6-100	1200-1218	FPKSLMTYANILLORRVPG	. 991	91	DR2-3-61	2268.6	2276.7
	177-100	NAGE I ASSISTANT 1130	167	. <b>9</b>	DR2-3-61	2127.4	2132.6
Potassius charmet proc		DGILYTYGSGGRLRRPV	168	11	DR2-3-61	2013.3	2018.1
fibronectin receptor	586-616	LSPIMIALWFSLDPQAPVDSHGLRPALHYQ	169	30	DR2-3-61	3307.7	3313.1
Factor VIII	1175-1790	LWDYGHSSSPHVLRWR	170	9 <b>:</b>	DR2-3-44	1918.2	1921.7
	07.111	#D JQD LXSQTVTVXQQVQ	2	51	0R2-3-39	2106.5	2114.
HLA-DKZO P-cnain	94-108	RVQPKVTVTPSKTQP	ĸ	15	DR2-3-39	1728.3	1730.6
							ESI-NS*
							27.0

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TABLE 5 MT-20/HLA-DR3 NATURALLY PROCESSED PEPTIDES

Protein source	Position	sednence	SEG 10 MO.	Length	Fraction	!	
Pseudo HIA-A2	103-117	VGSDWRFLRGYHOYA	2	15	DR3-2-63	1855.0	1863.9
WI A.A30	28.7	VDD1GFVRFDSDAASG	171	1	DR3-2-55	~	~
MIA-DO 4-chain	111-129	PPEVIVLINSPVELREPNV	71	19	DR3-2-55	50602	2093.3
	111-128	PPEVIVLINSPVELREPN	173	18	0R3-2-55	1991.2	1989.8
Ciedo. A Co. e In	1.2	COTRPRELEYSTSECHFF	٤	18	DR3-2-73	~	~
Academic for recent	289-104	VELLLADIXVPETSLS	174	<b>16</b>	DR3-2-65	1745.1	1750.1
Acetylonoline recept.	127-037	161614568066850	51	16	DR3-2-55	1670.8	1672.6
Glucose-transport	101-101	XXXIXXDIESUAEI	176	7	DR3-2-41	1720.8	1720.5
Sodium chambel prot.	204-377	diedmilgienskingsgedie	6	23	083-2-73	2545.2	2554.0
Invertent chain	08-110	DKDDKDVKWRWATPI I MOALP	. 24	25	DR3-2-73	2432.0	7.1772
<b>(E)</b>	90-110	KODKDVSKNRHATPLINGALP	5	12	DR3-2-73	2334.9	2345.3
	131-149	ATKYGWMTEDHVMHLLGWA	171	10	DR3-2-69	2173.4	2179.3
\$700	1071-1084	GQVKKNNHQEDKIE	178	7	DR3-2-41	1666.8	1667.0
ICAM-2	92-39	<b>LHKILLDEQAGUK</b>	179	13	DR3-2-51/52	1598.9	1602.4
	128-167	GPPKLDIRKEEKQIMIDIFH	180	21	DR3-2-77	2505.0	2510.3
Interreron 7-receptor		CODELDIBEEROIMIDIENP	181	20	DR3-2-77	2407.8	2412.4
•	02-02	SOLOALDEFENCEDOVINYETGNI	182	22	DR3-2-77	2505.0	2510.3
IP-30	18-57	SPI OAL DEFENGEPVNYKTG	251	20	083-2-77	2122.4	2124.2
Cytochrome-b5 reduc.	155-172	GKFAIRPOKKSNPIIRTV	181	18	083-2-51/52	2040.4	2043.2
EBV membrane antigen GP220	909-265	TGHGARTSTEPTTDY	185	<del>2</del>	DR3-2-41	1593.6	1592.7
EBV tegument protein membrane p140	1395-1407	KELKROYEKKIRO	<b>381</b>	žī.	DR3-2-51/52	. 1747.1	1749.8

Table 5, continue

	Position	Sequence	SEQ 10 NO.	Length	Frection	2	Nass Spec.
Apolipoprotein 1276	1276-1295	MFLKSDGRIKYTLMKNSLK	74	02	083-2-63	2352.9	2360.0
8-100 (Human) 1273	1273-1292	JPOML FLKSOGRIKYTLHKM	191	. 02	DR3-2-65	2349.7	2354.6
	1273-1291	I PDNL FLKSDGRIKYTLNK	ĸ	91	DR3-2-63	2235.5	. 1.242.1
127.	1273-1290	I PONL FLKSDGRJKYTLN	192	55	DR3-2-65	2107.4	2096.6
127	1273-1289	IPDNLFLKSDGRIKYTL	193	17	DR3-2-65	1993.3	2000.8
1274	1276-1291	HLFLKSDGRIKYTLNK	92	16	DR3-2-60	1910.2	1911.4
1276	1276-1290	NEFLKSDGRIKYTLM	11	15	DR3-2-60	1782.1	1785.9
1203	1207-1224	YANILLDRRVPQTONTF	78	17	DR3-2-63	2053.3	2059.1
¥.	1794-1810	VTTLNSDLKYNALDLTN	194	11	DR3-2-69	1895.1	1896.5
							MALD-MS

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TABLE &
PRIESS/HLA-DA4 MATURALLY PROCESSED PEPTIDES

PROTEIN SOURCE	P0S1710N	SEQUENCE	SEG 10 NO.	LENGTH	FRACTION	2	MASS SPEC
to Kampa Chain	168-208	KHKVYACEVTHOGLSSPVTKS .	80	12	DR4-2-45	2299.6	2304.0
range Cham	168-207	KHKVYACEVTHOGLSSPVTK	18	02	084-2-47	2212.5	2213.0
	189-206	HKVYACEVTHQGLSSPVT	82	18	DR4-2-43	1955.5	1952.1
	188-204	KHKYYACEVTWOGLSSP	83	17	084-2-45	1863.1	1882.8
	167-203	EKHKVYACEVTHOGLSS	Z	17	DR4-2-45	1915.1	1922.5
	188-203	KHKVYACEVTHOGLSS	88	16	DR4-2-54	1787.0	1787.0
	180-206	HKVYACEVTHOGLSSP	8	91	DR4-2-47	1755.0	1767.8
	167-202	EKHKVYACEVT HOGL S	87	16	DR4-2-43	1828.0	1822.8
	188-202	KHKVYACEVTHGGLS	8	15	084-2-51	1699.9	1708.3
	189-203	HKVYACEVTHOGLSS	89	<b>\$</b>	084-2-45	1657.8	1667.0
	187-200	EKHKVYACEVTHOG	8	7	DR4-2-51	1628.8	1632.6
	182.108	APSPLPETTERVYCALG	2	11	DR4-2-43	1697.9	1700
HLA-UK G-CASIII	28.50	VODITOEVREDSDAASORKEPRAP	195	23	084-2-58	2638.6	2641.5
74.45	87.82	VDDTOFVRFDSDAASQRHEPR	85	12	DR4-2-56	2470.6	2472.9
	28-47	VDDIGFYRFDSDAASGRMEP	8	50	DR4-2-59	2314.5	2319.3
	28.44	VODTOFVREDSDAASORKE	76	91	DR4-2-54	2217.2	2218.7
	87-UL	DIDEVAEDSDAASORMEPR	8	6	DR4-2-55	2256.4	2263.2
	67-11	TOFVREDSDAASGRMEPRA	96	61	084-2-56	2212.4	2211.5
	28-44	VDDTQFVRFDSDAASQR	26	11	DR4-2-55	1957.0	1963.1
	25-11	TOFVRFDSDAASGRMEP	86	11	DR4-2-56	1985.1	1987.5
	31-45	TOFVRFDSDAASORM	8	15	DR4-2-54	1758.9	1761.0
	67-F1	TOFVREDSDAAS	100	12	DR4-2-54	1343.4	1343.3
4	28-50	VDOTGEVREDSOAASPRGEPRAP		8	DR4-2-56	2533.7	2536.7
שרא_נ.	21-52	TOFVRFDSDAASPRGEPRAPLY	W 102	22	084-2-54	2,689.7	2491.5
	28-48	VDDTQFVRFDSOAASPAGEPR	103	12	DR4-2-54	2365.5	2368.1
	28-47	VODTOEVREDSOAASPRGEP	5	2	DR4-2-56	2509.3	2211.5
	;		;	;	73.67	2112 2	2111.0

cont inued
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	28-45	VODTOFVRFOSDAASPRG	106	18	DR4-2-56	1983.1	1987.5
חואינשי	W7-11	TOFVREDSDAASPRGEPR	101	5	0R4-2-52	2036.2	2041.5
	27 1C	VODTOFVREDSDAASPR	108	11	DR4-2-55	1926.0	1931.7
•	97-0£	DIDEVREDSDAASPRGE	109	11	DR4-2-52	1897.9	1901.6
	77-61	TOFVRFDSDAASPR	110	75	DR4-2-52	1596.7	1603.7
	27°F2	TOFVRFDSDAAS	=======================================	12	DR4-2-54	1343.4	1343.3
•	34-15 . USI-011	I RSWTAADTAAGTTORKWEAA	112	12	DR4-2-56	2374.6	7.9252
HLA.C	130-172	OI BOUTAADTAADTOBIN	197	4	DR4-2-58	2218.4	2220.1
	130-175	I RSUTANDIAAO I TORKW	198	18	DR4-2-58	2103.3	2105.0
	130-145	OI PSUTAADTAAOI TOR	113	11	DR4-2-59	1904.5	1908.7
	771-021	ni esutaditadi 10	116	91	DR4-2-59	1747.9	1752.3
	F/F-025	OLEGUTABOTAAQIT	115	5	084-2-59	1619.7	1622.2
	120-150	DI SSUTABDIAADITORKUEAA	<u>\$</u>	22	DR4-2-65	2420.6	2422.7
HLA-BWOZ	129-155	DI SSUTABOTABOTTOR	116	17	DR4-2-60	1834.9	1838.1
	771-061	DI SSVIAADTAAGITGRK	200	18	084-2-65	1963.1	1966.3
	129-148	DI SSUTAADTAAQTIQRICAE	117	20	DR4-2-66	2278.4	2284.6
•	970-066	GSLEVYNITTHKYKAFLDKO	201	20	DR4-2-65	2350.7	2352.6
VLA-4	776-066	CSE EVYNJ TINKYKAF	202	45	DR4-2-65	1866.1	1868.2
•	241-281	AADYFKFYPI SALTNILSAGL	203	2	084-2-65	2228.5	2229.5
PA1-1	107-107	AAPYFICEVPLSALTNILS	20%	18	DR4-2-65	1916.2	1917.4
	277-197	YDHNEVKATHADGKSVT	118	11	DR4-2-70	2037.2	2039.6
Cethepsin c			119			2035.3	
(Ret Momologue)	***	TO MODE TO A DOME OF	120	\$	084-2-70	1936.1	1937.7
	901-161					1934.2	
	17-76	AFALFRMFLSFPTIKT	202	16	DR4-2-78	1842.1	1836.1
Bovine Hemograpin	at-/c	SPEDEVAGECACY	506	15	084-2-78	1861.1	1.1981
HLA-043.2 p-chain	3 - 5	CDTRPRFLEQVKHE	122	7	084-2-72	1711.9	
IG Heavy Chain	121-7	GVYFYLOWGRSTLVSVS	123	3	DR4-2-6	~	-

TABLE 7
MANN/HLA-DR7 NATURALLY PROCESSED PEPTIDES

PRESENTED HILA-REZ         105-124         SOUNTELEGRINOTATOGGOTI         207         20         087-2-63           103-120         VICSDIAFI RGYNOTATOG         1         18         087-2-63           103-120         VICSDIAFI RGYNOTATOG         1         18         087-2-63           103-121         VICSDIAFI RGYNOTATOG         1         18         087-2-63           104-116         GSDUAFI RGYNOTATOG         2         15         087-2-63           105-117         SGDUAFI RGYNOTATOG         2         15         087-2-63           105-117         GSDUAFI RGYNOTATOG         209         13         087-2-63           105-117         SGDUAFI RGYNOTATOG         209         13         087-2-63           105-117         SGDUAFI RGYNOTATOG         12         187-2-63         087-2-63           105-117         SGDUAFI RGYNOTATOG         12         187-2-63         087-2-63           105-117         SGDUAFI RGYNOTATOG         12         187-2-63         087-2-63           110-128         RRADGI TGKUASVVVPSGG         12         187-2-63         087-2-63           110-128         RREDI SKYNITATOR         12         12         187-2-63           110-128         RREDI SKYNITATOR <t< th=""><th>PROTEIN SOURCE</th><th>POSITION</th><th>SEQUENCE</th><th>SEO 1D NO.</th><th>LENGTH</th><th>FRACT 10M</th><th>2</th><th>MASS SPEC</th></t<>	PROTEIN SOURCE	POSITION	SEQUENCE	SEO 1D NO.	LENGTH	FRACT 10M	2	MASS SPEC
103-120   VGSDMRFLRCTHOTATOC   1   18   103-120   VGSDMRFLRCTHOTATOC   1   18   104-115   VGSDMRFLRCTHOTA   2   15   104-116   GSDMRFLRCTHOTA   208   14   104-116   GSDMRFLRCTHOTA   209   13   105-117   SOURFLRCTHOTA   209   13   105-117   SOURFLRCTHOTA   209   13   105-117   SOURFLRCTHOTA   209   13   105-117   SOURFLRCTHOTA   209   13   105-125   GDGTFGKUASVVPSGGEGRYT   126   22   237-254   GDGTFGKUASVVPSGGEGRYT   126   22   239-225   GTGTAMSVVPSGGEGRYT   126   22   239-225   GTGTAMSVVPSGGEGRYTCHY   130   23   16   239-225   GTGTAMSVVPSGGEGRYTCHY   130   23   16   239-225   GTGTAMSVVPSGGEGRYTCHY   130   23   16   239-225   GTGTAMSVVPSGGGGRYTCHY   130   23   16   239-225   GTGTAMSVVPSGGGGRYTCHY   130   23   16   239-225   GTGTAMSVVPSGGGGRYTCHY   130   23   23   23   23   23   23   23	2	105.124	SOURFLEGYHOYAYDGKDY1	207	2	DR7-2-61	2553.8	2556.5
103-117   VIGSDMRLEGENOTA   2   15   16   16   17   16   17   16   17   18   18   18   19   19   19   19   19	Pseudo nia Ac	021 - CO1	SCYNORIE BOYNOVAYDG	-	8	DR7-2-63	2190.4	2194
103-117   CSDMFLEGTHOTA   208   14     104-116   CSDMFLEGTHOTA   209   13     105-117   SDMFLEGTHOTA   209   13     105-117   SDMFLEGTHOTA   209   13     105-117   SDMFLEGTHOTA   210   13     234-249   RPAGGGTGAMASVVPSGGE   124   20     237-254   GDGTGAMASVVPSGGE   127   16     239-253   GTFGAMASVVPSGGE   128   14     239-253   GTFGAMASVVPSGGE   129   15     239-261   GTFGAMASVVPSGGE   129   15     239-261   GTFGAMASVVPSGGE   129   15     239-263   GTFGAMASVVPSGGE   129   15     239-264   GTFGAMASVVPSGGE   129   15     239-265   GTFGAMASVVPSGGE   129   15     239-261   GTFGAMASVVPSGGGRYTCHV   130   23     239-263   GTFGAMASVVPSGGGRYTCHV   130   23     239-264   GTFGAMASVVPSGGGRYTCHV   130   23     239-265   GTFGAMASVVPSGGGRYTCHV   215   20     239-265   GTFGAMASVVPSGGGRYTCHV   215   21     24-200   APSPLPETTENVCALGLTV   215   21     24-200   APSPLPETTENVCALGLTV   215   21     24-200   APSPLPETTENVCALGT   230   14     25-200   KHKVYAGEVTHOGL   220   14     26-200   KHKVYAGEVTHOGL   220   14     26-200   KHKVYAGEVTHOGL   220   14     26-200   CAPHPRTUSGHLVGALFI   222   23     25-201   222   23     25-201   222   23     25-201   222   23     25-201   222   23     25-201   222   23     25-201   222   23     25-201   232   233     25-201   232     25-201   232     25-201   232     25-201   232     25-201   232		103-160	WACHAGE BORGON	~	<b>\$</b>	0R7-2-63	1855.0	1860
104-111 GSDMFLEGTHOT 209 13 105-117 SOURFLEGTHOT 209 13 105-117 SOURFLEGTHOT 209 13 234-259 RPAGGGTGGWASVVPSGG 124 20 234-269 RPAGGGTGGWASVVPSGGE 125 16 237-254 GDGTGGWASVVPSGGE 127 18 239-253 GTGGWASVVPSGGE 128 14 239-253 GTGGWASVVPSGGE 129 15 239-261 GTGGWASVVPSGGE 129 15 239-263 GTGGWASVVPSGGE 129 15 239-263 GTGGWASVVPSGG 129 15 239-264 GTGGWASVVPSGG 129 15 239-264 GTGGWASVVPSGG 129 15 239-265 GTGGWASVVPSGG 129 13 24-269 RETGISKTHIGTYREM 211 21 24-269 GALAMAVCANLEIMYGRAS 211 21 25-269 GTGGWASVVPSGGGRYTGHV 216 2 25-269 GTGGWASVVPSGGGRYTGHV 216 2 26-269 APSPLPETTENVCALGLTV 215 2 26-269 KKKVYAGEVTHQG 225 11 26-269 KKKVYAGEVTHQG 225 11 26-269 KKKVYAGEVTHQG 225 11 26-269 KKKVYAGEVTHQG 225 22 25-269 GONYPKTUSGHLWALP 13 21 26-269 GONYPKTUSGHLWALP 13 22 25 25-269 GONYPKTUSGHLWALLHAALP 13 22 25 25-269 GGGGWAS 10-10-10-10-10-10-10-10-10-10-10-10-10-1		103-111	TACHADA INGOOD	208	7	DR7-2-61	1755.9	1760.8
105-117 SOWRELRGTHOTA  234-253 RPAGGGTFGKMASVVPSGG 234-254 RPAGGGTFGKMASVVPSGG 234-259 RPAGGGTFGKMASVVPSGGGRYT 237-254 GGGTGKMASVVPSGGGRYT 237-254 GGGTGKMASVVPSGGGRYT 237-254 GGGTGKMASVVPSGGGRYT 239-255 GTGKMASVVPSGGGRYTCHY 239-255 GTGKMASVVPSGGGRYTCHY 239-255 GTGKMASVVPSGGGRYTCHY 239-256 GTGKMASVVPSGGGRYTCHY 239-266 GTGKMASVVPSGGGRYTCHY 239-267 RETGISKINIGTYREH 239-268 RETGISKINIGTYREH 239-269 RETGISKINIGTYREH 239-269 RETGISKINIGTYREH 239-261 GTGKMATOKANE HYKASH 239-261 GTGKMATOKANE HYKASH 231-375 GALAMANOKANE HYKASH 231-375 GALAMANOKANE HYKASH 231-375 GALAMANOKANE HYKASH 231-376 GA		104-117	AGHADA DONADO	50%	: 12	DR7-2-61	1684.8	1687.6
234-253         RPAGOGTFGKMASVVVPSGO         124         20           234-249         RPAGOGTFGKMASVVPSGQE ORT 1         125         16           237-254         GDGTFGKMASVVPSGQE         127         18           239-252         GTFGKMASVVPSGQE         127         18           239-253         GTFGKMASVVPSGQ         129         15           239-261         GTFGKMASVVPSGQ         129         15           239-263         GTFGKMASVVPSGQ         129         15           239-264         GTFGKMASVVPSGQ         129         15           239-265         GTFGKMASVVPSGQ         129         15           239-266         GTFGKMASVVPSGQ         129         15           239-267         GTFGKMASVVPSGQ         129         15           239-268         RETGISKINIGTREM         211         17           239-269         RETGISKINIGTREM         212         16           239-269         RETGISKINIGTREM         213         15           24-060         APSPLPETTENVCALGITV         214         26           250-76         GALAMIANDKASMISCIGGFV         216         17           eptor         179-7         SLGSP ITVENRAGSSGSAGSKMISCIGGFV         216		105-117	SOURFLEGYHOYA	210	13	DR7-2-61	1698.9	1704.1
24-259 RPAGDGTTGKMASVY 125 16 237-254 GDGTTGKMASVYPSGGEORTT 126 22 237-254 GDGTTGKMASVYPSGGEORTT 126 22 239-252 GDGTTGKMASVYPSGGE 127 18 239-253 GTGKMASVYPSGG 128 14 239-254 GDGTTGKMASVYPSGG 128 14 239-255 GTGKMASVYPSGG 129 15 239-261 GTGKMASVYPSGGGGRYTCHY 130 23 239-261 GTGKMASVYPSGGGGRYTCHY 130 23 239-261 GTGKMASVYPSGGGGRYTCHY 130 23 239-261 GTGKMASVYPSGGGGRYTCHY 130 23 239-261 GTGKMASVYPSGGGRYTCHY 130 23 239-261 GTGKMASVYPSGGGRYTCHY 130 23 239-261 GTGKMASVYPSGGGRYTCHY 130 23 24-259 GTGKMASVYPSGGGRYTCHY 130 23 24-259-259 GTGKMASVYPSGGRYTCHY 130 23 24-259 GTGKMASVYPSGGRANTPLHAALP 12 22 25 25-259 GTGKMASVYPSGRANTPLHAALP 13 22 25 25-259 GTGKMASVYPSGRANTPLHAALP 13 22 25			654 CANCEL SONIA CHANDERS	124	20	087-2-66	2087.3	2602
237-254 GDGTFQKUASVVPSGGEORTT 126 22 237-254 GDGTFQKUASVVPSGGE 127 18 239-252 GTFGKUASVVPSGGE 128 14 239-253 GTFGKUASVVPSGG 128 14 239-253 GTFGKUASVVPSGG 129 15 239-261 GTFGKUASVVPSGGEORTCHV 130 23 239-261 GTFGKUATGHTKRSH 211 21 239-261 GTFGKUATGHTKRSH 211 21 239-261 GTFGKUATGHTKRSH 211 21 239-261 GTFGKUATGHTKRSH 211 220 14 249-200 KHKYYACEVTHGG 220 14 249-200 KHKYYACEVTHGG 221 13 249-119 KPPRPVSKORRATPLLMQALP 13 21 249-119 KPPRPVSKORRATPLLMQALP 13 22 25	HLA-A29	60.40	O DACTO TECNIAL DO CONTRACTOR DE CONTRACTOR	125	\$	DR7-2-63	7171	1718
237-254 GOGTGGKAASVVPSGAE 127 18 239-255 GTFGKAASVVPSGAE 128 14 239-253 GTFGKAASVVPSGAE 129 15 239-254 GTFGKAASVVPSGAE 129 15 239-261 GTFGKASVVPSGAE 129 15 63-99 RETGISKTRIGTYRENL 211 17 63-99 RETGISKTRIGTYRENL 211 17 63-99 RETGISKTRIGTYREN 212 16 101-126 RSHYTPITNPFEVTVLTHSPVELREP 214 26 101-126 RSHYTPITNPFEVTVLTHSPVELREP 214 26 102-200 APSPLPETTEWVCALGLTV 215 20 179-7 SLGSPITVENRAGSESAGSKALSGIGGFVL 216 7 168-200 KHKYYACEVTHGGL 220 14 186-200 KHKYYACEVTHGGL 220 14 196-200 KHKYYACEVTHGGL 220 14 196-200 KHKYYACEVTHGGL 220 22 196-119 KPPPKPVSKORRATPLLMQALP 13 21 196-219 CAPAPACHUSALCALGALAGVLTI 222 25		497-967	TYBO BOS BONNESS CONTRACTOR TO THE CONTRACTOR TO	126	22	DR7-2-66	2436	0772
239-252 GTFGKUASVVPSG 128 14 239-253 GTFGKUASVVPSGQ 129 15 239-261 GTFGKUASVVPSGQ 129 15 239-261 GTFGKUASVVPSGQCGRYTCHV 130 23 239-261 GTFGKUASVVPSGQCGRYTCHV 130 23 63-99 RETGISKTHIGTYREN 211 17 63-96 RETGISKTHIGTYREN 2112 16 101-126 RSHYTPITNPPEVTVLTHSPVELREP 213 15 59-76 GALAMIAVDKAMLEIMTKRSM 131 21 59-76 GALAMIAVDKAMLEIMTKRSM 215 20 179-7 SLGSP ITVEURAGSESAGSKMLSGIGGFVL 216 7 179-179-179-179-179-179-179-179-179-179-		23-152	COGTFGKVASVVPSGGE	121	18	DR7-2-66	1892.3	1892
239-253 GTFGKVASVVPSGQ 129 15 239-261 GTFGKVASVVPPSGQEGRYTCHV 130 23 239-261 GTFGKVASVVPPSGQEGRYTCHV 130 23 63-99 RETGISKTHIGTYREHL 211 17 63-98 RETGISKTHIGTYREH 212 16 101-126 RSHYTP ITHPPEVTVLTHSPVELREP 214 26 101-126 RSHYTP ITHPPEVTVLTHSPVELREP 214 26 102-200 APSPLPETTENVCALGLTV 215 20 179-7 SLGSP ITVEURAGSESAGSKMLSGIGGFVL 216 7 179-7 SLGSP ITVEURAGSESAGSKMLSGIGGFVL 216 7 119-7 SLGSP ITVEURAGSESAGSKMLSGIGGFVL 216 7 119-336 VTGYLMATGHRUCSWSL 217 21 166-200 KHKYYACEVTHGGL 220 14 166-200 KHKYYACEVTHGGL 220 14 166-200 KHKYYACEVTHGGL 220 14 169-119 PKPPKPVSKMRMATPLLMAALP 12 22 96-119 RYPPKPVSKMRMATPLLMAALP 13 21 99-119 RYPPKPVSKMRMATPLLMAALP 13 22 22		230-252	GIFGKWASVVVPSG	128	75	087-2-66	1462	1465
239-261 GTFGKUASVVPSGGFGRYTCHV 130 23 63-99 RETGISKTHIGTYRENL 211 17 63-96 RETGISKTHIGTYRENL 212 16 101-126 RSHYTPITNPFEVTVLTHSPVELREP 214 26 101-126 RSHYTPITNPFEVTVLTHSPVELREP 214 26 102-200 APSPLPETTENVCALGLTV 215 20 179-7 SLGSPITVEVRAGSESAGSKALSGIGGFVL 216 7 168-200 KHKYYACEVTHGGL 220 14 168-200 KHKYYACEVTHGGL 220 14 198-201 KHKYYACEVTHGGL 220 14 198-201 KHKYYACEVTHGGL 220 14 198-201 KHKYYACEVTHGGL 220 22 199-119 KPPRPVSKORRATPLLMQALP 13 21 199-119 ZAPSKORRATPLLMQALP 222 25		230-253	GTFOKWASWWPSGO	129	15	DR7-2-66	1718	1221
63-99 RETGISKTHIGTTREHL 211 17 63-96 RETGISKTHIGTTREHL 212 16 63-97 RETGISKTHIGTTREH 212 16 101-126 RSHTPITHPFEVTVLTHSPVELREP 214 26 58-78 GALAMIAVOKANLEIMTKRSH 131 21 102-200 APSPLPETTENVVCALGITV 215 20 179-7 SLGSPITVENRAGSESAGSKNLSGIGGFVL 216 7 318-334 VTGYLMATGHRUCSWSLSGAR 217 21 654-866 ISILCTRKRENIK 219 13 654-860 KHKYYACEVTHGGL 220 14 186-200 KHKYYACEVTHGGL 220 14 196-200 KHKYYACEVTHGGL 220 14 196-200 CHRKYACEVTHGGL 220 14 196-200 CHRKYACEVTHGGL 220 14 196-200 CHRKYACEVTHGGL 220 22 96-119 KPPPKPVSKORRATPLLMQALP 13 21		239-261	GIFOKVASVVVPSGQEQRYICH	-	23	OR7-2-66	2603	9092
83-98   RETGISKTNIGTYREN   212   16     83-97   RETGISKTNIGTYRE   213   15     101-126   RSHYTPITHPPEVTVLTHSPVELREP   214   26     102-200   APSPLPETTENVYCALGITY   215   20     179-7   SLGSPITVENRAGSESAGSKALSGIGGFVL   216   2     179-7   SLGSPITVENRAGSESAGSKALSGIGGFVL   216   2     180-316   VIGTLMATGHRUCSMSL SGAR   217   2     180-201   KHKYYACEVTHGGL   220   14     180-200   KHKYYACEVTHGGL   220   14     180-200   KHKYYACEVTHGGL   220   14     180-219   PKPPKPVSKORRATPLLMGALP   12   22     190-119   KPPKPVSKORRATPLLMGALP   13   21     190-216   COMYPKTUSGALVGALCALAGVLTI   222   25     101-126   COMYPKTUSGALVGALCALAGVLTI   222   23     101-126   COMYPKTUSGALVGALCALAGVLTI   222   23     101-126   COMYPKTUSGALVGALCALAGVLTI   222   23     101-126   COMYPKTUSGALVGALCALAGVLTI   222   23     101-126   COMYPKTUSGALVGALCALAGALGALGALGALGALGALGALGALGALGALGALGALG	//0-4 ***	A1-00	RETOISKTHIGTYRENL		17	DR7-2-35	2082.3	2086.1
13	MLA-844	81.08	RETOLSKINIOTYREW	212	5	0R7-2-35	1969.1	1971.1
101-126   RSNTTP   INPPEVTVLTHSPVELREP   214   26     50-76		93.07	RETOISKINIOTYRE	213	25	DR7-2-35	1655.0	1857.3
18-76   SALAMIAYOKANLEINTKASH   131   21     18-76   APSPLPETTENVYCALGLIV   215   20     179-7   SLGSPITVENRAGSESAGSKALSGIGGFVL   216   7     318-336   VTGTLNATGNRUCSMSL SGAR   217   21     18-334   VTGTLNATGNRUCSWSL SGAR   218   17     654-866   TSILCYRKEUIK   218   17     188-201   KHKYYACEVTHGGL   220   14     188-200   KHKYYACEVTHGG   221   13     96-119   PKPPKPVSKORRATPLLMGALP   12   22     96-119   KPPKPVSKORRATPLLMGALP   13   21     09-119   KPPKPVSKORRATPLLMGALP   13   22     06-119   KPPKPVSKORRATPLLMGALP   13   21     06-119   KPPKPVSKORRATPLMGALP   13   21     06-119   KPRFPVSKORRATPLMGALP   13   21     06-119   KPPKPVSKORRATPLMGALP   13   21     06-119   MTGALP   14   25   25     06-119   MTGALP   15   25     06-119   MT		101-124	RSNYTPITAPPEVIVLINSPVELRE		92	DR7-2-35	2354.2	6.9262
102-200 APSPIPETTENVCALGLIV 215 20 179-7 SLOSPITTENVCALGLIV 215 20 318-336 VTQYLMATGNRAUCSMSLSGAR 217 21 318-334 VTQYLMATGNRUCSWSL 218 17 654-866 TSILCYRKREWIK 219 13 188-201 KHKYYACEVTHQGL 220 14 188-200 KHKYYACEVTHQGL 220 14 96-119 PKPPKPVSKORRATPLLMAALP 12 22 96-119 KPPKPVSKORRATPLLMAALP 12 22 70-516 GONYPKTUSGALVGALCALAGVLTI 222 25	HLA-DR a-chain	021-101	CALANTAVOKANI FIMIKESH		~	DR7-2-66	2229.5	1222
179-7   SLGSPITVEWRAGSESAGSKOLLSGIGGFVL 216		30-70	APSPI PETTENVCALGLIV	215	2	DR7-2-42	1912.2	1917.7
18-336   VIQTLNATGHRUCSWSLSGAR   217   21		170.7	SLOSPITVEWRAGSESAGSKALSGIC		~	DR7-2-35	~	2
13   13   13   14   15   17   18   17   18   17   18   18   17   18   18	ארע-מת מ-כוופונו	418.418	VIOYLWATCHRUCSUSESGAR		12	DR7-2-71	2.1245	2445.1
y chain 510-554  10 C reg. 188-201 KHKYTACEVTHGGL 220 14  10 Leg. 188-200 KHKYTACEVTHGG 221 13  ain 96-119 PKPPKPVSIORMATPLLMAALP 12 22  otein 492-516 GONYPKTUSGHLVGALCALAGVLTI 222 25	4F2 Cell-furrace	000-010	VIOVI MATCHBUTSUS!	210	11	DR7-2-71	1999.2	5001.9
In C reg. 188-201 KHKYYACEVTHGGL 220 14 188-200 KHKYYACEVTHGG 221 13 ain 98-119 PKPPKPVSIORMATPLLMOALP 12 22 otein 492-516 GONYPKTUSGMLVGALCALAGVLT1 222 25	antigen heavy chain	310-336	TOTAL CARROLL AND THE TOTAL AN	219	13	DR7-2-35	1696.0	1700.6
166-200 KHKYYACEYTHUG 96-119 PKPPKPVSIORNATPLLMAALP 12 22 99-119 KPPKPVSIORNATPLLMAALP 13 21 492-516 GOMYPKTUSGALVGALCALAGVLT1 222 25	Lif receptor	900-400		220	71	087-2-61	1612.9	1615.6
188-200 KHKVTALEVINAL 98-119 PKPPKPVSIORMATPLLMALP 12 22 99-119 KPPKPVSIORMATPLLMALP 13 21 492-516 GONYPKTUSGALVGALCALAGVLT1 222 25	1g keppe chain C reg.	102-891	NAME AND ADDRESS OF THE PARTY O	<u> </u>	: =	DR7-2-61	1498.7	1501.0
96-119 PKPPKPUSKNKMAIPLLMAALP 13 21 99-119 KPPKPUSKNKMAIPLLMAALP 13 21 492-516 GONYPKTUSGNLVGALCALAGVLTI 222 25		186-200	KAKATACKATAGE	: :	: %	087-2-72	2432.0	2436.6
99-119 NFFAFFSMANDAIFLEMANT 222 25 492-516 CONTPETUDIONI VGALCALAGOLTI 222 25	Invariant Chain	98-119	PAPPAPASANAMITELMANT	: #	: 5	DR7-2-72	23%.9	2339.7
492-516 COMYPKTUSCALVGALCALAGVLII 222 23	3	9-119	KPRFVSKARDAIPLLMALF		: %	np7-2-71	2567.1	2567.3
	K channel protein	492-516	COMYPKTUSCHLVGALCALAGVLTI		Q	7.140		

Table 7, continued

PROTEIN SOURCE	POS1110#	SEQUENCE	SEQ 10 110.	LENGTH	FRACTION	Z	MASS SPEC
	18.57	IPSTVAFTOTERL ICDA	132	17	0R7-2-69	1856.0	1856.6
Heat Shock Cognate	K R			11	DR7-2-72	1856.0	1857.0
/1 KD protein	10.63	TPSYVAFTOTERLIG	133	5	DR7-2-69	1669.8	1671.9
	37-05	ADVI 1805 SPOT INDA	223	61	087-2-61	2079.5	2083.9
Complement Ly Thromboxane-A	707-750	PAFRFTREAAGOCEV	526	51	DR7-2-71	1739.9	1743.0
synthase	0801-7761	VOCI VODERAFIERE	525	81	0R7-2-54	2082.4	2.1902
EBV mejor cepsid prot	2021 - 1765	VOCI VSPCBAFFUK	226	7	DR7-2-54	1597.9	1598.6
	1721:-4071	KYDLIST CHALL CSDYGADYES	227	23	DR7-2-54	5660.9	5.2992
Apol ipoprotein 8-100	1500-1600	CODI TESCONAL LCS	228	15	DR7-2-54	1689.0	1,7891
	1700-1054	ESHOYBOSTSHB1	622	13	DR7-2-42	1562.7	1567.5
	2077-2089	LPKYFEKKRNTII	230	13	DR7-2-61	1650.0	1653.8
							MALD-MS

TABLE 8
Z3.1/MLA-DR8 MATURALLY PROCESSED PEPTIDES

PROTETH SOURCE	POSITION	SEGLENCE	SEQ. 10 NO.	LENGTH	FRACTION	2	MSS SPEC
		O 150 INDUSTRIBUTION CONTRACTOR	231	S	DR8-3-59	2889.3	2889.0
KLA-DR a-chain	158-160	ADEDI DETTENAÇÃI G	232	11	DR8-3-41	1697.9	1704.3
	182-198		213	~	DR8-3-75	•	•
HLA-DR B-chain	1-7	GOINFRILETS! UCC. 17 TOLEN	717	2	088-3-76	1587.7	1591.3
HLA-DP B-choin	80-92	RHWYELDEAVILE	3,5	: 2	DR8-3-54	2543.6	2549.1
LAM Blast-1 with	88-108	DPGSGALYI SKVQKEDNSI TI	246	17	DR8-3-52	2116.1	2118.0
M-acetyglucosamine	92-108	CALTISKVAKEONSITI	217	. 52	DR8-3-57	2081.4	2085.7
	129-146	DPVPKPVIKIEKIEURUU	238	15	DR8-3-57	1720.0	1724.9
	129-143	DPVPKPVIKIEKIEU	010	81	0R8-3-57	2201.5	2203.6
lg kappe chain	63-80	FIFTISKLEPEDIAVIVE	157	55	0R8-3-57	1772.0	1777.0
	63-77	FIFTISALEPEOPAV	24.2	. 71	DR8-3-76	1675.9	1679.8
LAR protein	1302-1316	OPVEMRRLNYOTPG	773	2 5	DRB-3-66	2108.5	2112.0
LIF receptor	924-604	YOLLRSMIGYTEELAPIV	2,3	2 2	088-3-66	2072.4	2075.1
IFN-a receptor	271-287	GHHLYKVKGIPOCENVK	3	: 6	08-1-80	2400.7	2402.5
Interleukin-6	169-188	LPFFLFRGATHPNNSSPVCY	544	2			
receptor	1	340X00AAQOYJATGGGGAAQOXAA	572	28	DR8-2-63	3161.6	3164.9
Metalloproteinase	187-214	QAKFFACIKKSDUSCAMIKUMATRACE	Ì				
Inhibitor 2			<b>97</b> 6		DRB-3-63	2235.5	2233.6
	187-205	CAKFFACIKKSUGSLANTK		φ.	DR8-3-66	2040.3	2042.9
Metalloproteinase	101-118	NRSEEFL I AGKL QDGL L H	<u>*</u>	2			
inhibitor 1			116	16	DR8-3-70	1789.0	1799.9
	101-117	· SEEPLIAGELY	276	15	DR8-3-72	1632.9	1646.0
	103-117	SEEFLIAGRICUGEL	378	2	088-3-66	1376.6	1381.8
	101-112	ARSEEFLIAGKL	0 × C	. 72	DRB-3-59	2662.9	3.4992
Cathepsin E	89-112	ONFTVIFOTGSSMLWPSVTC1SF	<b>(5</b> )	: 2	088-3-63	1857.9	1857.1
Cathepsin S	189-205	TAFOYIONKGIOSDAS	8 5	: 2	088-3-63	2348.7	2348.0
Cystelin SN	41-58	DETYRRILRYLRAREGIV	65 53	: 4	0R8-3-63	2077.3	2078.3
Tubuiln a-1 chain	207-223	EAITOICARNLOIEAPI	5 9	: =	088-3-63	1593.8	1595.1
	207-219	EALYOICRRMLDI	252	5 52	DR8-3-59	2493.9	0.4645
Myosin A-heavy chain	1027-1047	HELEKIKKOVEGEKCETGAAL PPSMI OHILIR	3 2	<u>.</u> 0	DRB-3-66	1250.5	1254.8
Ca release charrel	707-7102	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1					

Table 8, continued

PROTEIN SOURCE	P051710#	SEGLENCE	SEQ. 10 NO.	LENGTH	FRACTION	3	MSS SPEC
		AT ID INDIAN COURSE	255	22	DR8-3-72	2417.8	2421.3
C035	359-380	UNITED TO STATE OF THE STATE OF	ĸ	17	DRB-3-66	2195.6	1.2023
27.2	106-122	IPHLUKIMANILSONA	× ×	7	DR8-3-68	1706.0	1709.6
c-myc transfor. prot.	371-385	KASTALKULITOL		17	DR8-3-54	5064.4	2066.5
K-res tresnfor, prot. Celcitonin	164-180 38-53	EPFLYILGKSRVLEAQ	69	91	DR8-3-78	1863.2	1878.4
receptor (Hum?)			250	~	DR8-3-54	:	:
a-ENOLASE (7)	23-7	AEVYHDVAASE FF	5 52	. 4	0R8-3-59	2246.7	1.7255
Plasminogen activator	378-396	DRPFLFVVRHNPSGIVLFM	3, 3,	91	DRG-3-70	2008.4	2116.4
inhibitor-1	133-148	MPHF FRL FRST VKGVO	197	? 5	DR8-3-62	2393.8	2399.4
Apolipoprotein 8-100	1724-1743	KNI FHFKVNGEGLKI SNDRM	. 297	; <del>*</del>	DRB-3-57	1902.2	1903.7
	1724-1739	KMIFHFKVNDEGLKLS	<b>503</b>	2 02	DRB-3-54	2271.5	2273.7
	1780-1799	YKOTVSLOIOPYSLVITLMS	<b>6</b>	2 2	088-3-60	1918.2	1929.4
	2646-2662	STPEFTILNTLMIPSFT	607	. 15	0RB-3-60	2059.3	2073.5
	2647-2664	TPEFTICHTCHIPSFILD	8 2	. 91	DR8-3-80	1831.1	1841.6
	2647-2662	TPEFTICNICAIPSFT	103	: 5	DR8-3-68	1965.2	1969.9
	2885-2900	SHTKYFHKINIPOLDF	88 56	21	OR8-3-75	2203.6	2207.0
	2072-2088	LPFFKFLPKYFEKKRNT	607	: ¥	DR8-3-76	1988.4	1992.6
	2072-2086 4022-4036	LPFFKFLPKYFEKKR UMFYYSPQSSPDKKL	S €	21	DR8-3-59	1860.0	1863.3
			22	12	DR8-3-76	523.8	2524.9
Bovine Transferrin	261-281	DVINE LLXRAGENT GROKSKE	. K	51	DR8-3-78	1508.0	1818.1
	261-275 261-273	DVIWELLIMHAGENTG BVIVELLMHAGEH	196	ī.	DR8-3-73	1603.8	1608.8
		Sylvan and a sylva	061	20	DR6-3-59	2360.8	2359.7
von Willebrand factor	617-636	IALLIMASGEPORM	169	7	0R8-3-59	1600.9	1601.3

TABLE 9 HONZ/HIA-DR1 MATURALLY PROCESSED PEPTIDES

Pseudo HLA-A2   103-117   VGSDNRFLRGTHOTA	PROTEIN SOURCE	POSITION	SEGUENCE	SEQ 10 NO.	LENGTH	FRACTION	2	MASS SPEC
97-120         LPKPPKPVSKNRMATPLLMOALPHG         7         24         HZ/DR1-1-77         2676.4           98-121         PKPPKPVSKNRMATPLLMOALPHG         8         24         HZ/DR1-1-72         2620.2           97-119         LPKPPKPVSKNRMATPLLMOALPH         9         23         HZ/DR1-1-73         2545.2           96-120         PKPPKPVSKNRMATPLLMOALPH         10         23         HZ/DR1-1-75         2563.2           99-120         KPPKPVSKNRMATPLLMOALPH         11         22         HZ/DR1-1-75         2432.0           99-119         PKPPKPVSKNRMATPLLMOALPH         13         21         HZ/DR1-1-72         2334.9	Pseudo HLA-A2	103-117	VGSDURFLRGTHOTA GSDURFLRGTHOTA	2 2	15	N2/DR1-1-64 N2/DR1-1-63	1855.0	1854.4 1755.2
	Invertent Chain (11)	97-120 98-121 97-119 98-120 99-120 96-119	LPKPPKPVSKHRMATPLLMDALPHG PKPPKPVSKHRMATPLLMDALPHG LPKPPKPVSKHRMATPLLMDALP PKPPKPVSKHRMATPLLMDALPH RPPKPVSKHRMATPLLMDALPH KPPKPVSKHRMATPLLMDALP	7 8 9 11 11 12 13	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	HZ/DR1-1-77 HZ/DR1-1-72 HZ/DR1-1-73 HZ/DR1-1-75 HZ/DR1-1-75 HZ/DR1-1-72	2676.4 2620.2 2545.2 2563.2 2466.1 2432.0	2619.7 2619.7 2544.5 2562.3 2455.8 2431.7 2334.2

SUMMARY OF MATURALLY PROCESSED PEPTIDES BOUND TO HIA-DR EXPRESSED IN MORMAL HUMAN SPLEEN

HIA-DR a-chain 71/133-156 SETVFLPREDHLFRKFHYLPFLPS 140 24 2976 2665  71/136-156 VFLPREDHLFRKFHYLPFLPS 141 20 20 2577 2579  71/136-151 VFLPREDHLFRKFHYLPFLPS 142 20 2577 2579  71/136-153 KLGHPDILM 144 99 994 994 999  Calgranulin B 33/25-33 KLGHPDILM MASHEKHHEGDEGFRHHHKPGLGEGTP 145 277 2017 2017 2017  42/80-114 WASHEKHHEGDEGFRHHHKPGLGEGTP 146 27 2017 2017 2023  KLA-B51 42/104-121 GPDGRLLRGHHWPGLGEGTP 146 1704 1704 1704 1705  KINASE C f chain (rat) 42/341-446 ILPPFGPGINOYGLD 70 16 16 1704 1899 1894  HIA-DR # chain 45/129-144 WASHERHGGEKTOWS 71 16 16 1892 1894	PROTEIN SOURCE	POS1110#	SEQUENCE	SEO 10 NO.	LENGTH	3	MASS SPEC	
71/136-156 VFLPREDHLFRKFHYLPFLP 142 20 2572 71/136-155 VFLPREDHLFRKFHYLPFLP 142 20 2572 71/136-151 VFLPREDHLFRKFHYLPFLP 143 16 2118 33/25-33 KLGHPDILM 42/88-114 WASHEKHHEGDEGPGHHKPGLGEGTP 145 27 2017 42/88-114 WASHEKHHEGDEGPGHHKPGLGEGTP 146 27 2017 in (ret) 42/341-446 ILPPFGPGITDDYGLD 70 16 1704  n 45/129-144 VRUFRNGGEEKTGVVS 71 16 1892	HA-08 archain	71/133-156	SETVFLPREDHLFRKFHYLPFLPS	140	*	2976	2982	
71/136-155 VFLPREDHLFRKFHYLPFLP 142 20 2572 71/136-151 VFLPREDHLFRKFHYLPFLP 143 16 2118 33/25-33 KLGHPDTLM 42/88-114 UASHEKHHEGDEGPGHHKFGLGEGTP 145 27 2915 43/88-114 UASHEKHHEGDEGPGHHKFGLGEGTP 146 27 2915 42/88-114 UASHEKHHEGDEGPGHHKFGLGEGTP 146 27 2017 42/104-121 GPOGRLLRGHWYDGK 188 16 2017 42/341-446 TLPPFGPG1TDDYGLO 70 16 1704 45/129-144 VRUFRNGGEEKTGVVS 71 16 1892		71/136-156	VFLPREDHLFRKFHYLPFLPS	171	12	592	5992	
71/136-151         VFLPREDHLFRKFHYL         143         16         2118           33/25-33         KLGHPDTLM         144         9         994           42/88-114         WASHEKNHEGDEGPGHHKPGLGEGTP         145         27         2915           43/88-114         WASHEKNHEGDEGPGHHKPGLGEGTP         146         27         2017           42/104-121         GPOGRLLRGHNOYDGK         188         16         2017           42/341-446         TLPPFGPQ1TDDYGLD         70         16         1704           45/129-144         VRUFRNGGEEKTGVVS         71         16         1704		71/136-155	VFLPREDKLFRKFHYLPFLP	142	20	2572	25.79	
33/25-33 KLGHPDTLM 144 9 994 42/88-114 WASHERHHEGDEGPCHHHKPGLGEGTP 145 27 2915 43/88-114 WASHERHHEGDEGPCHHHKPGLGEGTP 146 27 2017 42/104-121 GPOGRLLRGHWGYDGK 188 16 2017 42/341-446 TLPPFGPQ1TDDYGLO 70 16 1704 45/129-144 VRUFRNGGEEKTGVVS 71 16 1892		71/136-151	VFLPREDKLFRKFHYL	143	<b>3</b> 1	2118	2126	
42/68-114         WASHEKHHEGDEGPCHHHKPGLGEGTP         145         27         2915           43/68-114         WASHEKHHEGDEGPCHHHKPGLGEGTP         146         27         2017           42/104-121         GPOGRLLRGHWOYDGK         188         16         2017           42/341-446         TLPPFGPG1TDDYGLD         70         16         1704           45/129-144         VRUFRNGGEEKTGVVS         71         16         1892	. B ci juganojan	33/25-33	KIGHPOTIN	144	۰	%	666	
43/88-114 WASHERHHEGDEGPGHHKPGLGEGTP 146 27 2017 42/104-121 GPOGRLLRGHWOYDGK 188 16 2017 42/341-446 TLPPFGPQ1TDDYGLO 70 16 1704 45/129-144 VRWFRNGGEEKTGVVS 71 16 1892		42788-114	MASHEKNHEGDEGPGHHHKPGLGEGTP	145	27	2915	1262	
42/104-121 GPOGRLLRGHWOYDGK 188 16 2017 42/341-446 TLPPFGPGITODYGLD 70 16 1704 45/129-144 VRWFRNGGEEKTGVVS 71 16 1892		43/88-114	WASHEKMHEGDEGPGHHHKPGLGEGTP	971	22	2017	5926	
42/341-446 TLPPFAPAITODYGLO 70 16 1704 45/129-144 VRUFRNGAEEKTGVVS 71 16 1892	HLA-851	42/104-121	GPOGRLLRGHNOYDGK	188	16	2017	2023	
45/129-144 VRUFRNGGEEKTGVVS 71 16 1892	Kinase C f chain (rat)	75/341-446	11 <i>PPF</i>	02	. 91	1704	1705	
SH-gTN	HLA-DR6 ß chain	45/129-144	VRUFRNGGEEKTGVVS	2	92	1892	1894	
							MALD-MS	

PCT/US93/07545

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## SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT:

Robert G. Urban Roman M. Chicz Dario A. A. Vignali Mary L. Hedley Lawrence J. Stern Jack L. Strominger

(ii) TITLE OF INVENTION:

IMMUNOMODULATORY PEPTIDES

(iii) NUMBER OF SEQUENCES:

273

(iv) CORRESPONDENCE ADDRESS:

(A) ADDRESSEE:

Fish & Richardson 225 Franklin Street

(B) STREET: (C) CITY:

Boston

(D) STATE:

Massachusetts

(E) COUNTRY:

U.S.A.

(F) ZIP:

02110-2804

(v) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: 3.5" Diskette, 1.44 Mb
(B) COMPUTER: IBM PS/2 Model 50Z or 55SX
(C) OPERATING SYSTEM: MS-DOS (Version 5.0)

(D) SOFTWARE:

WordPerfect (Version 5.1)

(vi) CURRENT APPLICATION DATA:

(A) APPLICATION NUMBER:

(B) FILING DATE:

(C) CLASSIFICATION:

(vii) PRIOR APPLICATION DATA:

(A) APPLICATION NUMBER: 07/925,460

(B) FILING DATE:

August 11, 1992

(viii) ATTORNEY/AGENT INFORMATION:

Clark, Paul T. 30,162

(A) NAME: (B) REGISTRATION NUMBER:

(C) REFERENCE/DOCKET NUMBER: 00246/168001

5:	3 -
(ix) TELECOMMUNICATION INF	ORMATION:
	17) 542-5070 17) 542-8906 0154
(2) INFORMATION FOR SEQUENCE IDENTI	FICATION NUMBER: 1:
(i) SEQUENCE CHARACTERISTI	cs:
(A) LENGTH: 18 (B) TYPE: amino ac (C) STRANDEDNESS: (D) TOPOLOGY: 1:	rid .near
(xi) SEQUENCE DESCRIPTION:	SEQ ID NO: 1:
Val Gly Ser Asp Trp Arg Phe Leu Arg	G Gly Tyr His Gln Tyr Ala Tyr 10 15
Asp Gly	
(2) INFORMATION FOR SEQUENCE IDENT	
(i) SEQUENCE CHARACTERIST	ics:
(A) LENGTH: 15 (B) TYPE: amino a (C) STRANDEDNESS: (D) TOPOLOGY: 1	cid inear
(xi) SEQUENCE DESCRIPTION	: SEQ ID NO: 2:
Val Gly Ser Asp Trp Arg Phe Leu Ar 1 5	g Gly Tyr His Gln Tyr Ala 10 15
	· _

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (1) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 14
      (B) TYPE: amino acid
      (C) STRANDEDNESS:
      (D) TOPOLOGY: linear linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Val Gly Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr 1

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 14
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

- 54 -

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

Gly Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 13
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 25
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

Leu Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro

Leu Leu Met Gln Ala Leu Pro Met Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (1) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 24
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

Leu Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro 1 5 10 15

Leu Leu Met Gln Ala Leu Pro Met 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 24
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu 1 5 10

Leu Met Gln Ala Leu Pro Met Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

Leu Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro

Leu Leu Met Gln Ala Leu Pro 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 23 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu

Leu Met Gln Ala Leu Pro Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu Leu

Met Gln Ala Leu Pro Met

- .... (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu

Leu Met Gln Ala Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 13:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu Leu

Met Gln Ala Leu Pro 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid

(C) STRANDEDNESS:

linear (D) TOPOLOGY:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu Leu Met

Gln Ala Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

Lys Met Arg Met Ala Thr Pro Leu Leu Met Gln Ala Leu Pro Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 16:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - linear (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

Lys Met Arg Met Ala Thr Pro Leu Leu Met Gln Ala Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 17:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - amino acid (B) TYPE:

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

Ile Pro Ala Asp Leu Arg Ile Ile Ser Ala Asn Gly Cys Lys Val Asp

Asn Ser

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

Arg Val Glu Tyr His Phe Leu Ser Pro Tyr Val Ser Pro Lys Glu Ser

Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

Tyr Lys His Thr Leu Asn Gln Ile Asp Ser Val Lys Val Trp Pro Arg

Arg Pro Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 20:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

Tyr Lys His Thr Leu Asn Gln Ile Asp Ser Val Lys Val Trp Pro Arg

Arg Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 21:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 19
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - linear (D) TOPOLOGY:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

Asp Val Gly Glu Tyr Arg Ala Val Thr Glu Leu Gly Arg Pro Asp Ala 1 5 10 15

Glu Tyr Trp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 22:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

Glu Pro Gly Glu Pro Glu Phe Lys Tyr Ile Gly Asn Met His Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 17 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

Tyr Lys His Thr Leu Asn Gln Ile Asp Ser Val Lys Val Trp Pro Arg

Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 24:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

Pro Lys Tyr Val Lys Gln Asn Thr Leu Lys Leu Ala Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15.
    - (B) TYPE: amino acid

- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

Leu Pro Lys Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr 1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 26:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

Ser Asp Leu Ser Phe Ser Lys Asp Trp Ser Phe Tyr Leu
10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 27:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

Lys Val Phe Gly Arg Cys Glu Leu Ala Ala Ala Met Lys Arg His Gly 1 5 10 15

Leu Asp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 28:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

Arg Asn Arg Cys Lys Gly Thr Asp Val Gln Ala Trp Ile Arg Gly Cys 10 15

Arg Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 29:
  - (i) SEQUENCE CHARACTERISTICS:

(A) LE	NGTH:	16
--------	-------	----

- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY:

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

His Pro Pro His Ile Glu Ile Gln Met Leu Lys Asn Gly Lys Lys Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 16
      (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

Asn Glu Leu Gly Arg Phe Lys His Thr Asp Ala Cys Cys Arg Thr His

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

Ser Lys Pro Lys Val Tyr Gln Trp Phe Asp Leu Arg Lys Tyr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32:

Ala Thr Ser Thr Lys Lys Leu His Lys Glu Pro Ala Thr Leu Ile Lys

Ala Ile Asp Gly 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 20
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

Pro Ala Thr Leu Ile Lys Ala Ile Asp Gly Asp Thr Val Lys Leu Met
1 5 10 15

Tyr Lys Gly Gln 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 34:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

Asp Arg Val Lys Leu Met Tyr Lys Gly Gln Pro Met Thr Phe Arg Leu

Leu Leu Val Asp 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 35:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 35:

Val Ala Tyr Val Tyr Lys Pro Asn Asn Thr His Glu Gln His Leu Arg

Lys Ser Glu Ala 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 36:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 36:

Gln Lys Gln Glu Pro Ile Asp Lys Glu Leu Tyr Pro Leu Thr Ser Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 37:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 16 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

Gly Ala Arg Ala Ser Val Leu Ser Gly Gly Glu Leu Asp Lys Trp Glu
1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 38:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 38:

Arg Thr Leu Tyr Gln Asn Val Gly Thr Tyr Val Ser Val Gly Thr Ser

Thr Leu Asn Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 39:

Leu Lys Lys Leu Val Phe Gly Tyr Arg Lys Pro Leu Asp Asn Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:

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(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:

.Lys His Ile Glu Gln Tyr Leu Lys Lys Ile Lys Asn Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:

Asp Val Phe Lys Glu Leu Lys Val His His Ala Asn Glu Asn Ile Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 42:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 42:

Gly Asp Thr Arg Pro Arg Phe Leu Trp Gln Leu Lys Phe Glu Cys His

Phe Phe Asn Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 22 (B) TYPE: am amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 43:

Thr Glu Arg Val Arg Leu Leu Glu Arg Cys Ile Tyr Asn Gln Glu Glu 1 10 15

Ser Val Arg Phe Asp Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 44:
  - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTE: 25
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- linear (D) TOPOLOGY:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 44:

Asp Leu Leu Glu Gln Arg Arg Ala Ala Val Asp Thr Tyr Cys Arg His

Asn Tyr Gly Val Gly Glu Ser Phe Thr -20 25

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 45:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:

Lys Ala Glu Arg Ala Asp Leu Ile Ala Tyr Leu Lys Gln Ala Thr Ala 1 10 15

Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 24
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:

Gly Arg Thr Gln Asp Glu Asn Pro Val Val His Phe Phe Lys Asn Ile

Val Thr Pro Arg Thr Pro Pro Pro 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 47:

Pro Leu Lys Ala Glu Ile Ala Gln Arg Leu Glu Asp Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 48:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48:

Arg Gln Ile Leu Gly Gln Leu Gln Pro Ser Leu Gln Thr Gly Ser Glu
1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 49:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 49:

Ile Gln Val Tyr Ser Arg His Pro Pro Glu Asn Gly Lys Pro Asn Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 50: .
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 16 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 50:

Ile Asn Thr Lys Cys Tyr Lys Leu Glu His Pro Val Thr Gly Cys Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 51:

Tyr Lys Leu Asn Phe Tyr Phe Asp Leu Leu Arg Ala Lys Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 52:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 52:

Ile Asp Thr Leu Lys Lys Asn Glu Asn Ile Lys Glu Leu
1 5 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 53:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 20 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 53:

Asp Val Gly Glu Tyr Arg Ala Val Thr Glu Leu Gly Arg Pro Asp Ala

Glu Tyr Trp Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 16 (B) TYPE: am amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 54:

Glu Arg Phe Ala Val Asn Pro Gly Leu Leu Glu Thr Ser Glu Gly Cys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid

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- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 55:

Asp Asn Tyr Arg Gly Tyr Ser Leu Gly Asn Trp Val Cys Ala Ala Lys

Phe Glu Ser Asn Phe Thr Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 56:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 56:

Glu Ala Leu Val Arg Gln Gly Leu Ala Lys Val Ala Tyr Val Tyr Lys

Pro Asn Asn Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPB: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 57:

Pro Ile Val Gln Asn Leu Gln Gly Gln Met Val His Gln Ala Ile Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 58:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS:
      (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 58:

Ser Ala Leu Ser Glu Gly Ala Thr Pro Gln Asp Leu Asn Thr Met Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 59:
  - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 16
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 59:

Ser Phe Tyr Ile Leu Ala His Thr Glu Phe Thr Pro Thr Glu Thr Asp
1 5 10 15

- (2) IMPORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 60:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 60:

Lys Met Tyr Phe Asn Leu Ile Asn Thr Lys Cys Tyr Lys Leu Glu His 1 5 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 61:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 61:

Val Gly Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala Tyr
1 5 10 15

Ala Asp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 62:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 62:

Gly Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala Tyr Asp 1 5 10 15

Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 63:
  - (i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 20
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 63:

Pro Pro Lys Pro Val Ser Lys Met Arg Met Ala Thr Pro Leu Leu Met

Gln Ala Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 64:

Lys Met Arg Met Ala Thr Pro Leu Leu Met Gln Ala Leu Pro Met Gly

Ala Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 65:

Glu His His Ile Phe Leu Gly Ala Thr Asn Tyr Ile Tyr Val Leu Asn

Glu Glu Asp Leu Gln Lys Val 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 66:

Gln Glu Leu Lys Asn Lys Tyr Tyr Gln Val Pro Arg Lys Gly Ile Gln

Ala

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 67:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 67:

Ile Gln Asn Leu Ile Lys Glu Glu Ala Phe Leu Gly Ile Thr Asp Glu
1 5 10 15

Lys Thr Glu Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 68:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 68:

Thr Ala Phe Gln Tyr Ile Ile Asp Asn Lys Gly Ile Asp Ser Asp Ala 1 5 10 15

Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 69:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 69:

Glu Pro Phe Leu Tyr Ile Leu Gly Lys Ser Arg Val Leu Glu Ala Gln
1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 70:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 70:

Thr Leu Pro Pro Phe Gln Pro Gln Ile Thr Asp Asp Tyr Gly Leu Asp 1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 71:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 71:

Val Arg Trp Phe Arg Asn Gly Gln Glu Glu Lys Thr Gly Val Val Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 72:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 72:

Arg Val Gln Pro Lys Val Thr Val Tyr Pro Ser Lys Thr Gln Pro Leu 1 5 10 15

Gln His

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 73:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 73:

Arg Val Gln Pro Lys Val Thr Val Tyr Pro Ser Lys Thr Gln Pro 1 5 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 74:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:

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- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 74:

Asn Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr Leu Asn Lys Asn 1 5 10

Ser Leu Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 19
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 75:

Ile Pro Asp Asn Leu Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr 1 5 10 15

Leu Asn Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 16
  (B) TYPE: \_ amino acid

  - (C) STRANDEDNESS: (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 76:

Asn Leu Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr Leu Asn Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 77:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 77:

Asn Leu Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr Leu Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 17

(B) TYPE: amino acid

(C) STRANDEDNESS:

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 78:

Tyr Ala Asn Ile Leu Leu Asp Arg Arg Val Pro Gln Thr Asp Met Thr

Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 79:

Gly Asp Thr Arg Pro Arg Phe Leu Glu Tyr Ser Thr Ser Glu Cys His

Phe Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 80:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 80:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser 1 5 10

Pro Val Thr Lys Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 81:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser

Pro Val Thr Lys

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 82:

His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser Pro

Val Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 83:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 84:

Glu Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser

Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 85:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 16
    - amino acid (B) TYPE:

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 86:

His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 87:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - amino acid (B) TYPE:
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 87:

Glu Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - amino acid (B) TYPE:

    - (C) STRANDEDNESS: (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 88:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 89:

His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 90:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 90:

Glu Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 91:

Ala Pro Ser Pro Leu Pro Glu Thr Thr Glu Asn Val Val Cys Ala Leu

Glý

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 21
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 92:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln

Arg Met Glu Pro Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 20
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

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(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 93:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln

Arg Met Glu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 94:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln 10 15

Arg Met Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 95:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (x1) SEQUENCE DESCRIPTION: SEQ ID NO: 95:

Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln Arg Met

Glu Pro Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 19

    - (B) TYPE: amino acid
      (C) STRANDEDNESS:
      (D) TOPOLOGY: lines linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 96:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln Arg Met Glu

Pro Arg Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 97:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 97:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln

Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 98:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln Arg Met Glu

Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 99:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln Arg Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

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(A) LENGTE: 12

(B) TYPE: amino acid

(C) STRANDEDNESS:

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 100:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser 1

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 101:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 23 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 101:

Val Asp Asp Thr Gin Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro

Arg Gly Glu Pro Arg Ala Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 102:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 102:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro Arg Gly Glu

Pro Arg Ala Pro Trp Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 21
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: (D) TOPOLOGY:
    - linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 103:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro

Arg Gly Glu Pro Arg 20

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 104:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 104:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro

Arg Gly Glu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 105:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 105:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro

Arg Gly Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 106:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 106:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro

Arg Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 18
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 107:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro Arg Gly Glu 10

Pro Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 108:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 108:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro

Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 109:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 109:

Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro Arg Gly

Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 110:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Pro Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 12

PCT/US93/07545

- amino acid (B) TYPE:
- (C) STRANDEDNESS:
- (D) TOPOLOGY:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 111:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 112:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 112:

Leu Arg Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln Arg

Lys Trp Glu Ala Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 17
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 113:

Asp Leu Arg Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln
10

Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 114:

Asp Leu Arg Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 115:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 15
- (B) TYPE: amino acid (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 115:

Asp Leu Arg Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 116:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 116:

Asp Leu Ser Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln 1 5 10 15

Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 117:

Asp Leu Ser Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln 1 5

Arg Lys Trp Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 118:

Tyr Asp His Asn Phe Val Lys Ala Ile Asn Ala Asp Gln Lys Ser Trp

Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 119:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 119:

Tyr Asp His Asn Phe Val Lys Ala Ile Asn Ala Asp Ile Lys Ser Trp

Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - .. (A) LENGTH: 16
      - (B) TYPE: amino acid (C) STRANDEDNESS:

      - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 120:

Tyr Asp His Asn Phe Val Lys Ala Ile Asn Ala Asp Gln Lys Ser Trp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 121:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 121:

Tyr Asp His Asn Phe Val Lys Ala Ile Asn Ala Ile Gln Lys Ser Trp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 122:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:

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- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 122:

Gly Asp Thr Arg Pro Arg Phe Leu Glu Gln Val Lys His Glu
1 5 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 123:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 123:

Gly Val Tyr Phe Tyr Leu Gln Trp Gly Arg Ser Thr Leu Val Ser Val
1 5 10 15

Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 124:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 124:

Arg Pro Ala Gly Asp Gly Thr Phe Gln Lys Trp Ala Ser Val Val Val 1 1 15

Pro Ser Gly Gln 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 125:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 125:

Arg Pro Ala Gly Asp Gly Thr Phe Gln Lys Trp Ala Ser Val Val 11 15

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 126:

- (i) SEQUENCE CHARACTERISTICS:

  - (A) LENGTE: 22 (B) TYPE: amino acid
  - (C) STRANDEDNESS:
  - (D) TOPOLOGY:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 126:

Gly Asp Gly Thr Phe Gln Lys Trp Ala Ser Val Val Val Pro Ser Gly

Gln Glu Gln Arg Tyr Thr 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 18 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 127:

Gly Asp Gly Thr Phe Gln Lys Trp Ala Ser Val Val Val Pro Ser Gly
1 5 10 15

Gln Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 128:

Gly Thr Phe Gln Lys Trp Ala Ser Val Val Val Pro Ser Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 129:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - amino acid (B) TYPE:
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 129:

Gly Thr Phe Gln Lys Trp Ala Ser Val Val Val Pro Ser Gly Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 130:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 130:

Gly Thr Phe Gln Lys Trp Ala Ser Val Val Val Pro Ser Gly Gln Glu

Gln Arg Tyr Thr Cys His Val 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21
    - (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 131:

Gly Ala Leu Ala Asn Ile Ala Val Asp Lys Ala Asn Leu Glu Ile Met
1 5 10 15

Thr Lys Arg Ser Asn 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - amino acid (B) TYPE:
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 132:

Thr Pro Ser Tyr Val Ala Phe Thr Asp Thr Glu Arg Leu Ile Gly Asp

Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 133:

Thr Pro Ser Tyr Val Ala Phe Thr Asp Thr Glu Arg Leu Ile Gly
1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 134:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 134:

Arg Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu Gln Asp Gly Leu Leu
1 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 135:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 135:

Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu Gln Asp Gly Leu Leu
1 5 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 136:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 136:

Asp Val Ile Trp Glu Leu Leu Asn His Ala Gln Glu His Phe Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 137:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 137:

Glu Pro Phe Leu Tyr Ile Leu Gly Lys Ser Arg Val Leu Glu Ala Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 138:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 15
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 138:

Thr Ala Phe Gln Tyr Ile Ile Asp Asn Lys Gly Ile Asp Ser Asp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 139:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 139:

Thr Ala Phe Gln Tyr Ile Ile Asp Asn Lys Gly Ile Asp Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 140:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 24
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 140:

Ser Glu Thr Val Phe Leu Pro Arg Glu Asp His Leu Phe Arg Lys Phe 10

His Tyr Leu Pro Phe Leu Pro Ser

- (2) IMPORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 141:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 21 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 141:

Val Phe Leu Pro Arg Glu Asp His Leu Phe Arg Lys Phe His Tyr Leu
1 10 15

Pro Phe Leu Pro Ser 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 142:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 142:

Val Phe Leu Pro Arg Glu Asp His Leu Phe Arg Lys Phe His Tyr Leu
1 10 15

Pro Phe Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 143:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 143:
- Val Phe Leu Pro Arg Glu Asp His Leu Phe Arg Lys Phe His Tyr Leu 1 10 15
- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 144: (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 26
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS:
  - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 144:

Lys Leu Gly His Pro Asp Thr Leu Asn Gln Gly Glu Phe Lys Glu Leu 1 5 15

· Val Arg Lys Asp Leu Gln Asn Phe Leu Lys 20 25

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 145:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 24

ATG GCC ATA Met Ala Ile

CTG ATG AGC Leu Met Ser

CTG CTG ATG Leu Leu Met 35

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(B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 145:	
Lys Leu Gly His Pro Asp Thr Leu Asn Gln Gly Glu Phe Lys Glu Leu 1 5 10	•
Val Arg Lys Asp Leu Gln Asn Phe 20	
(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 146:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTE: 14 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 146:	
Lys Leu Gly His Pro Asp Thr Leu Asn Gln Gly Glu Phe Lys 1 5 10	
(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 147:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 123 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 147:	
G GCC ATA AGT GGA GTC CCT GTG CTA GGA TTT TTC ATC ATA GCT GTG  t Ala Ile Ser Gly Val Pro Val Leu Gly Phe Phe Ile Ile Ala Val  5 10 15	48
G ATG AGC GCT CAG GAA TCA TGG GCT AAG ATG CGC ATG GCC ACC CCG Bu Met Ser Ala Gln Glu Ser Trp Ala Lys Met Arg Met Ala Thr Pro 20 25 30	96
ng CTG ATG CAG GCG CTG CCC ATG TAA nu Leu Met Gln Ala Leu Pro Met 35 40	123
) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 148:	
(i) SEQUENCE CHARACTERISTICS:	

- (2) INFORMAT
  - - (A) LENGTH: 150
      (B) TYPE: nucleic acid
      (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear

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## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 148:

- ATG GCC ATA AGT GGA GTC CCT GTG CTA GGA TTT TTC ATC ATA GCT GTG Met Ala Ile Ser Gly Val Pro Val Leu Gly Phe Phe Ile Ile Ala Val
- CTG ATG AGC GCT CAG GAA TCA TGG GCT CTT CCC AAG CCT CCC AAG CCT 96 Leu Met Ser Ala Gln Glu Ser Trp Ala Leu Pro Lys Pro Pro Lys Pro
- GTG AGC AAG ATG CGC ATG GCC ACC CCG CTG CTG ATG CAG GCG CTG CCC Val Ser Lys Met Arg Met Ala Thr Pro Leu Leu Met Gln Ala Leu Pro 40

150 ATG TAA Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 10
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 149:

Thr Gln Phe Val Arg Phe Asp Ser Asp Ala . 1

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 150:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 150:

Asp Trp Arg Phe Leu Arg Gly Tyr His Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 151:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 151:

Arg Met Ala Thr Pro Leu Leu Met Gln Ala

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(B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 152: Lys Asp Glu Leu (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 5 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 153: Lys Phe Glu Arg Gln (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: (i) "SEQUENCE CHARACTERISTICS: (A) LENGTE: 5
(B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 154: Gln Arg Glu Phe Lys (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 155: Met Ala Ile Ser Gly Val Pro Val Leu Gly Phe Phe Ile Ile Ala Val Leu Met Ser Ala Gln Glu Ser Trp Ala

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 156:

Met Arg Met Ala Thr Pro Leu Leu Met Gln Ala Leu Pro Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 157:

Met Pro Arg Ser Arg Ala Leu Ile Leu Gly Val Leu Ala Leu Thr Thr 1 5 10 15

Met Leu Ser Leu Cys Gly Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 158:

Asn Ile Val Ile Lys Arg Ser Asn Ser Thr Ala Ala Thr Asn Glu Val

Pro Glu Val Thr Val Phe Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY:

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 159:

Asn Ile Val Ile Lys Arg Ser Asn Ser Thr Ala Ala Thr Asn Glu Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 18 (B) TYPE: amino acid (C) STRANDEDNESS:

    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 160:

Ser Asp Val Gly Val Tyr Arg Ala Val Thr Pro Gln Gly Arg Pro Asp

Ala Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 161:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17

    - (B) TYPE: amino acid
      (C) STRANDEDNESS:
      (D) TOPOLOGY: line
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 161:

Asp Val Gly Val Tyr Arg Ala Val Thr Pro Gln Gly Arg Pro Asp Ala 1 5 10 15

Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 162:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 162:

Asp Val Gly Val Tyr Arg Ala Val Thr Pro Gln Gly Arg Pro Asp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 163:
  - (i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 13
- (B) TYPE: amino acid (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 163:

Ala Pro Ser Pro Leu Pro Glu Thr Thr Glu Asn Val Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 164:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 17 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 164:

Ala Pro Ser Pro Leu Pro Glu Thr Thr Glu Asn Val Val Cys Ala Leu

Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 165:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 165:

Phe Pro Lys Ser Leu His Thr Tyr Ala Asn Ile Leu Leu Asp Arg Arg

Val Pro Gln Thr Asp 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 166:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 166:

Phe Pro Lys Ser Leu His Thr Tyr Ala Asn Ile Leu Leu Asp Arg Arg

Val Pro Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 167:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 167:

Asp Gly Ile Leu Tyr Tyr Gln Ser Gly Gly Arg Leu Arg Arg Pro 1 5 10

Val Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 168:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 168:

Asp Gly Ile Leu Tyr Tyr Gln Ser Gly Gly Arg Leu Arg Arg Pro

Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 30
    - amino acid
    - (B) TYPE: amino (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 169:

Leu Ser Pro Ile His Ile Ala Leu Asn Phe Ser Leu Asp Pro Gln Ala

Pro Val Asp Ser His Gly Leu Arg Pro Ala Leu His Tyr Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 170:

Leu Trp Asp Tyr Gly Met Ser Ser Pro His Val Leu Arg Asn Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 171:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 16 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 171:
- Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln
- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 172:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 19

    - (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: line:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 172:

Pro Pro Glu Val Thr Val Leu Thr Asn Ser Pro Val Glu Leu Arg Glu

Pro Asn Val

- (2) INFORMATION POR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 173:

Pro Pro Glu Val Thr Val Leu Thr Asn Ser Pro Val Glu Leu Arg Glu

Pro Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 174:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 174:

Val Phe Leu Leu Leu Ala Asp Lys Val Pro Glu Thr Ser Leu Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 175:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 175:

Thr Phe Asp Glu Ile Ala Ser Gly Phe Arg Gln Gly Gly Ala Ser Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 176:

Tyr Gly Tyr Thr Ser Tyr Asp Thr Phe Ser Trp Ala Phe Leu
1 5 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid (C) STRANDEDNESS:

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(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 177:

Ala Thr Lys Tyr Gly Asn Met Thr Glu Asp His Val Met His Leu Leu

Gln Asn Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 178:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 14
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 178:
- Gly Gln Val Lys Lys Asn Asn His Gln Glu Asp Lys Ile Glu 1 10
- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 179:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 179:

Leu Asn Lys Ile Leu Leu Asp Glu Gln Ala Gln Trp Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 180:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 180:

Gly Pro Pro Lys Leu Asp Ile Arg Lys Glu Glu Lys Gln Ile Met Ile

Asp Ile Phe His

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 181: - 102 -

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 181:

Gly Pro Pro Lys Leu Asp Ile Arg Lys Glu Glu Lys Gln Ile Met Ile

Asp Ile Phe His Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 182:

Ser Pro Leu Gln Ala Leu Asp Phe Phe Gly Asn Gly Pro Pro Val Asn

Tyr Lys Thr Gly Asn Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 183:

Ser Pro Leu Gln Ala Leu Asp Phe Phe Gly Asn Gly Pro Pro Val Asn 1 10 15

Tyr Lys Thr Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 184:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 184:

Gly Lys Phe Ala Ile Arg Pro Asp Lys Lys Ser Asn Pro Ile Ile Arg

Thr Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 185:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 15
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 185:

Thr Gly His Gly Ala Arg Thr Ser Thr Glu Pro Thr Thr Asp Tyr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 186:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 13 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 186:

Lys Glu Leu Lys Arg Gln Tyr Glu Lys Lys Leu Arg Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 187:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 12
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 187:

Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 188:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 16 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 188:

Gly Pro Asp Gly Arg Leu Leu Arg Gly His Asn Gln Tyr Asp Gly Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 189:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 14
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 189:

Ile Ala Leu Leu Met Ala Ser Gln Glu Pro Gln Arg Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 190:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 190:

Ile Ala Leu Leu Met Ala Ser Gln Glu Pro Gln Arg Met Ser Arg

Asn Phe Val Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 20
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 191:

Ile Pro Asp Asn Leu Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr

Leu Asn Lys Asn

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTE: 18
  - (B) TYPE: amino acid (C) STRANDEDNESS:

  - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 192:

Ile Pro Asp Asn Leu Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr

Leu Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 193:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 193:

Ile Pro Asp Asn Leu Phe Leu Lys Ser Asp Gly Arg Ile Lys Tyr Thr

Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 17 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 194:

Val Thr Thr Leu Asn Ser Asp Leu Lys Tyr Asn Ala Leu Asp Leu Thr 1 10 15

Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 195:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 195:

Val Asp Asp Thr Gln Phe Val Arg Phe Asp Ser Asp Ala Ala Ser Gln

Arg Met Glu Pro Arg Ala Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 196:

Asp Val Ile Trp Glu Leu Leu Asn His Ala Gln Glu His

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 197:

Asp Leu Arg Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln

Arg Lys Trp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 198:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 18 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 198:

Leu Arg Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln Arg

Lys Trp

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 199:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 22
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 199:

Asp Leu Ser Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln 1 5 10 15

Arg Lys Trp Glu Ala Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 200:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 200:

Asp Leu Ser Ser Trp Thr Ala Ala Asp Thr Ala Ala Gln Ile Thr Gln

Arg Lys

- 201: (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 20 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 201:

Gly Ser Leu Phe Val Tyr Asn Ile Thr Thr Asn Lys Tyr Lys Ala Phe

Leu Asp Lys Gln

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 202:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid

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- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 202:

Gly Ser Leu Phe Val Tyr Asn Ile Thr Thr Asn Lys Tyr Lys Ala Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 21 (B) TYPE: amino acid

    - (C) STRANDEDNESS:
      (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 203:

Ala Ala Pro Tyr Glu Lys Glu Val Pro Leu Ser Ala Leu Thr Asn Ile 10 .

Leu Ser Ala Gln Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 204:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
      - (B) TYPE: amino acid (C) STRANDEDNESS:

      - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 204:

Ala Ala Pro Tyr Glu Lys Glu Val Pro Leu Ser Ala Leu Thr Asn Ile

Leu Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 205:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 205:

Ala Glu Ala Leu Glu Arg Met Phe Leu Ser Phe Pro Thr Thr Lys Thr

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 206:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 206:

Ser Pro Glu Asp Phe Val Tyr Gln Phe Lys Gly Met Cys Tyr Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 207:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 207:

Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala Tyr Asp Gly

Lys Asp Tyr Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 208:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 208:

Gly Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala 1 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 209:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 209:

Gly Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 210:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - amino acid (B) TYPE:
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 210:

Ser Asp Trp Arg Phe Leu Arg Gly Tyr His Gln Tyr Ala

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 211:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 211:

Arg Glu Thr Gln Ile Ser Lys Thr Asn Thr Gln Thr Tyr Arg Glu Asn

Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 212:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 212:

Arg Glu Thr Gln Ile Ser Lys Thr Asn Thr Gln Thr Tyr Arg Glu Asn

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 213:
  - (1) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 15
    - (B) TYPE: amino acid

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- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 213:

Arg Glu Thr Gln Ile Ser Lys Thr Asn Thr Gln Thr Tyr Arg Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 26
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 214:

Arg Ser Asn Tyr Thr Pro Ile Thr Asn Pro Pro Glu Val Thr Val Leu

Thr Asn Ser Pro Val Glu Leu Arg Glu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 215:

Ala Pro Ser Pro Leu Pro Glu Thr Thr Glu Asn Val Val Cys Ala Leu

Gly Leu Thr Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 216:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 216:

Ser Leu Gln Ser Pro Ile Thr Val Glu Trp Arg Ala Gln Ser Glu Ser 1 5 10 15

Ala Gln Ser Lys Met Leu Ser Gly Ile Gly Gly Phe Val Leu

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- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 217:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 217:

Val Thr Gln Tyr Leu Asn Ala Thr Gly Asn Arg Trp Cys Ser Trp Ser

Leu Ser Gln Ala Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 218:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - amino acid
    - (B) TYPE: amino (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 218:

Val Thr Gln Tyr Leu Asn Ala Thr Gly Asn Arg Trp Cys Ser Trp Ser

Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 219:

Thr Ser Ile Leu Cys Tyr Arg Lys Arg Glu Trp Ile Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 220:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 220:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 221:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 13 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 221:

Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 222:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 25
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 222:

Gly Asp Met Tyr Pro Lys Thr Trp Ser Gly Met Leu Val Gly Ala Leu

Cys Ala Leu Ala Gly Val Leu Thr Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 223:

Ala Pro Val Leu Ile Ser Gln Lys Leu Ser Pro Ile Tyr Asn Leu Val

Pro Val Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15

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- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 224:

Pro Ala Phe Arg Phe Thr Arg Glu Ala Ala Gln Asp Cys Glu Val 1 5 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 225:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 225:

Val Pro Gly Leu Tyr Ser Pro Cys Arg Ala Phe Phe Asn Lys Glu Glu
1 10 15

Leu Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 226:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 226:

Val Pro Gly Leu Tyr Ser Pro Cys Arg Ala Phe Phe Asn Lys 1 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 227:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 227:

·Lys Val Asp Leu Thr Phe Ser Lys Gln His Ala Leu Leu Cys Ser Asp 1 5 15

Tyr Gln Ala Asp Tyr Glu Ser

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(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 15
- (B) TYPE: amino acid (C) STRANDEDNESS:
- (D) TOPOLOGY:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 228:

Lys Val Asp Leu Thr Phe Ser Lys Gln His Ala Leu Leu Cys Ser

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 229:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - linear (D) TOPOLOGY:
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 229:

Phe Ser His Asp Tyr Arg Gly Ser Thr Ser His Arg Leu
1 2 10

- 230: (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 13
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 230:

Leu Pro Lys Tyr Phe Glu Lys Lys Arg Asn Thr Ile Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 231:

Ser Glu Thr Val Phe Leu Pro Arg Glu Asp His Leu Phe Arg Lys Phe

His Tyr Leu Pro Phe Leu Pro

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 18
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 232:

Ala Pro Ser Pro Leu Pro Glu Glu Thr Thr Glu Asn Val Val Cys Ala Leu Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 233:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 24
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 233:

Gly Asp Thr Arg Pro Arg Phe Leu Glu Tyr Ser Thr Gly Glu Cys Tyr 1 5 10 15

Phe Phe Asn Gly Thr Glu Arg Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 13 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 234:

Arg His Asn Tyr Glu Leu Asp Glu Ala Val Thr Leu Gln

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 235: - 117 -

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTE: 21
  - (B) TYPE: amino acid (C) STRANDEDNESS:

  - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 235:

Asp Pro Gln Ser Gly Ala Leu Tyr Ile Ser Lys Val Gln Lys Glu Asp

Asn Ser Thr Tyr Ile 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 236:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 236:

Gly Ala Leu Tyr Ile Ser Lys Val Gln Lys Glu Asp Asn Ser Thr Tyr

Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 237:

Asp Pro Val Pro Lys Pro Val Ile Lys Ile Glu Lys Ile Glu Asp Met 10 15

Asp Asp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 238:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 15
      (B) TYPE: amino acid
      (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 238:

Asp Pro Val Pro Lys Pro Val Ile Lys Ile Glu Lys Ile Glu Asp 1 5 10 15

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 239:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 239:

Phe Thr Phe Thr Ile Ser Arg Leu Glu Pro Glu Asp Phe Ala Val Tyr

Tyr Cys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 240:

Phe Thr Phe Thr Ile Ser Arg Leu Glu Pro Glu Asp Phe Ala Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 241:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 241:

Asp Pro Val Glu Met Arg Arg Leu Asn Tyr Gln Thr Pro Gly

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 242:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid

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- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 242:

Tyr Gln Leu Leu Arg Ser Met Ile Gly Tyr Ile Glu Glu Leu Ala Pro 1 5 15

Ile Val

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 243:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 243:

Gly Asn His Leu Tyr Lys Trp Lys Gln Ile Pro Asp Cys Glu Asn Val

Lys

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 244:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 244:

Leu Pro Phe Phe Leu Phe Arg Gln Ala Tyr His Pro Asn Asn Ser Ser 1 5 10 15

Pro Val Cys Tyr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 245:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 28
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 245:

Gln Ala Lys Phe Phe Ala Cys Ile Lys Arg Ser Asp Gly Ser Cys Ala

Trp Tyr Arg Gly Ala Ala Pro Pro Lys Gln Glu Phe 20 25

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 19
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 246:

Gln Ala Lys Phe Phe Ala Cys Ile Lys Arg Ser Asp Gly Ser Cys Ala 1 10 15

Trp Tyr Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 15
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 247:

Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu Gln Asp Gly Leu Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 12 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 248:

Asn Arg Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu 1 5 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 24
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- linear (D) TOPOLOGY:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 249:

Gln Asn Phe Thr Val Ile Phe Asp Thr Gly Ser Ser Asn Leu Trp Val

Pro Ser Val Tyr Cys Thr Ser Pro 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 250:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 250:

Asp Glu Tyr Tyr Arg Arg Leu Leu Arg Val Leu Arg Ala Arg Glu Gln

Ile Val

- (2) IMPORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 251:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 251:

Glu Ala Ile Tyr Asp Ile Cys Arg Arg Asn Leu Asp Ile Glu Arg Pro

Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 252:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTE: 13 (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 252:

Glu Ala Ile Tyr Asp Ile Cys Arg Arg Asn Leu Asp Ile

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 253:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 21
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 253:

His Glu Leu Glu Lys Ile Lys Lys Gln Val Glu Gln Glu Lys Cys Glu

Ile Gln Ala Ala Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 254:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 10
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 254:

Arg Pro Ser Met Leu Gln His Leu Leu Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 255:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 255:

Asp Asp Phe Met Gly Gln Leu Leu Asn Gly Arg Val Leu Phe Pro Val

Asn Leu Gln Leu Gly Ala 20

- . (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 256:

Ile Pro Arg Leu Gln Lys Ile Trp Lys Asn Tyr Leu Ser Met Asn Lys

Tyr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 14
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 257:

Lys Arg Ser Phe Phe Ala Leu Arg Asp Gln Ile Pro Asp Leu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 258:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 258:

Arg Gln Tyr Arg Leu Lys Lys Ile Ser Lys Glu Glu Lys Thr Pro Gly 1 5

Сув

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 259:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 259:

Ala Glu Val Tyr His Asp Val Ala Ala Ser Glu Phe Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 260:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 19

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- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 260:

Asp Arg Pro Phe Leu Phe Val Val Arg His Asn Pro Thr Gly Thr Val

Leu Phe Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 261:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid

    - (C) STRANDEDNESS:
      (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 261:

Met Pro His Phe Phe Arg Leu Phe Arg Ser Thr Val Lys Gln Val Asp

- (2) IMPORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 262:

Lys Asn Ile Phe His Phe Lys Val Asn Gln Glu Gly Leu Lys Leu Ser

Asn Asp Met Met

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 16
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 263:

Lys Asn Ile Phe His Phe Lys Val Asn Gln Glu Gly Leu Lys Leu Ser

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 264: **-** 125 -

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 264:

Tyr Lys Gln Thr Val Ser Leu Asp Ile Gln Pro Tyr Ser Leu Val Thr

Thr Leu Asn Ser 20

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 17
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 265:

Ser Thr Pro Glu Phe Thr Ile Leu Asn Thr Leu His Ile Pro Ser Phe

Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 18
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 266:

Thr Pro Glu Phe Thr Ile Leu Asn Thr Leu His Ile Pro Ser Phe Thr 5 10 1

Ile Asp

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 267:

Thr Pro Glu Phe Thr Ile Leu Asn Thr Leu His Ile Pro Ser Phe Thr 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 268:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - linear (D) TOPOLOGY:
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 268:

Ser Asn Thr Lys Tyr Phe His Lys Leu Asn Ile Pro Gln Leu Asp Phe

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17
    - (B) TYPE: amino acid (C) STRANDEDNESS:

    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 269:

Leu Pro Phe Phe Lys Phe Leu Pro Lys Tyr Phe Glu Lys Lys Arg Asn

Thr

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 270:

Leu Pro Phe Phe Lys Phe Leu Pro Lys Tyr Phe Glu Lys Lys Arg

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:
  - (1) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 15

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- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 271:

Trp Asn Phe Tyr Tyr Ser Pro Gln Ser Ser Pro Asp Lys Leu
1 5 10

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 272:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 21
      (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 272:

Asp Val Ile Trp Glu Leu Leu Asn His Ala Gln Glu His Phe Gly Lys

Asp Lys Ser Lys Glu

- (2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 273:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTE: 16
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 273:

Asp Val Ile Trp Glu Leu Leu Ile Asn His Ala Gln Glu His Phe Gly

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## CLAIMS

- 1. A purified preparation of a peptide consisting essentially of an amino acid sequence identical to that of a segment of a naturally-occurring human protein, said segment being of 10 to 30 residues in length, inclusive, wherein said peptide binds to a human major histocompatibility complex (MHC) class II allotype.
- 2. The preparation of claim 1, wherein said peptide binds to at least two distinct MHC class II allotypes.
- The preparation of claim 1, wherein said human protein is HLA-A2, HLA-A29, HLA-A30, HLA-B44, HLA-B51, HLA-Bw62, HLA-C, HLA-DP  $\beta$ -chain, HLA-DQ  $\alpha$ -chain, HLA-DQ  $\beta$ chain, HLA-DQ3.2  $\beta$ -chain, HLA-DR  $\alpha$ -chain, HLA-DR  $\beta$ -chain, HLA-DR4  $\beta$ -chain, invariant chain (Ii), Ig kappa chain, Ig kappa chain C region, Ig heavy chain, Na+/K+ ATPase, potassium channel protein, sodium channel protein, calcium release channel protein, complement C9, glucose-transport protein, CD35, CD45, CD75, vinculin, calgranulin B, kinase C  $\zeta$ -chain, integrin  $\beta$ -4 gp150, hemoglobin, tubulin  $\alpha$ -1 myosin  $\beta$ -heavy chain,  $\alpha$ -enolase, transferrin, transferrin receptor, fibronectin receptor acetylcholine receptor, interleukin-8 receptor, interferon  $\alpha$ -receptor, interferon  $\gamma$ -receptor, calcitonin receptor, LAM (lymphocyte activation marker) Blast-1, LAR (leukocyte antigen-related) protein, LIF (leukemia inhibitory factor) receptor, 4F2 cell-surface antigen (a cell-surface antigen involved in normal and neoplastic growth) heavy chain, cystatin SN, VLA-4 (a cell surface heterodimer in the adhesion receptors), PAI-1 superfamily of integrin (plasminogen activator inhibitor-1), IP-30 (interferon-y induced protein), ICAM-2, carboxypeptidase E, thromboxane-A synthase, NADH-cytochrome-b5 reductase, c-myc transforming protein, K-ras transforming protein, MET kinase-related

transforming protein, interferon-induced guanylate-binding protein, mannose-binding protein, apolipoprotein B-100, cathepsin C, cathepsin E, cathepsin S, Factor VIII, von Willebrand factor, metalloproteinase inhibitor 1 precursor, metalloproteinase inhibitor 2, plasminogen activator inhibitor-1, or heat shock cognate 71 kD protein.

- 4. The preparation of claim 1, wherein said human protein is an MHC class I or II molecule.
- 5. The preparation of claim 1, wherein said segment conforms to the following motif:

at a first reference position (I) at or within 12 residues of the amino terminal residue of said segment, a positively charged residue or a large hydrophobic residue; and

at position I+5, a hydrogen bond donor residue.

- 6. The preparation of claim 5, wherein said motif comprises a hydrophobic residue at I+9.
- 7. The preparation of claim 6, wherein said motif additionally comprises, at position I+1 or I-1, a hydrophobic residue.
- 8. The preparation of claim 1, wherein said segment comprises residues 29-40 (SEQ ID NO: 187) or residues 106-115 (SEQ ID NO: 150) of HLA-A2.
- 9. The preparation of claim 1, wherein said segment comprises residues 107-116 of Ii (SEQ ID NO: 151).

- 10. A liposome containing a peptide consisting essentially of an amino acid sequence identical to that of a segment of a naturally-occurring human protein, said segment being of 10 to 30 residues in length, wherein said peptide binds to a human major histocompatibility complex (MHC) class II allotype.
- 11. An immune-stimulating complex (ISCOM) comprising a peptide consisting essentially of an amino acid sequence identical to that of a segment of a naturally-occurring human protein, said segment being of 10 to 30 residues in length, wherein said peptide binds to a human major histocompatibility complex (MHC) class II allotype.
- 12. A nucleic acid encoding a polypeptide, said polypeptide comprising a first and a second amino acid sequence linked by a peptide bond, said first sequence being identical to that of a segment of a naturally-occurring human protein, which segment binds to a human MHC class II allotype and is of 10 to 30 residues in length; and said second sequence being a sequence which controls intracellular trafficking of a polypeptide to which it is attached ("trafficking sequence").
- 13. The nucleic acid of claim 12, wherein said trafficking sequence is KDEL (SEQ ID NO: 152); KFERQ (SEQ ID NO: 153); QREFK (SEQ ID NO: 154); MAISGVPVLGFFIIAVLMSAQESWA (SEQ ID NO: 155); a pentapeptide comprising Q flanked on one side by four residues selected from K, R, D, E, F, I, V, and L; or a signal peptide.
- 14. A nucleic acid encoding a polypeptide comprising a first and a second amino acid sequence linked by a peptide bond, said first sequence being identical to that

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of a segment of a naturally-occurring human protein, which segment binds to a human MHC class II allotype and is of 10 to 30 residues in length; and said second sequence being substantially identical to human Ii.

- 15. A cell comprising the nucleic acid molecule of claim 14.
  - 16. A method of making a peptide, which method comprises culturing the cell of claim 15 under conditions permitting expression of said peptide from said nucleic acid molecule.
  - 17. The preparation of claim 1, wherein said segment consists essentially of a sequence set forth in any of Tables 1-10.
  - 18. A method of identifying a nonallelically restricted immunomodulating peptide, said method comprising:
  - (a) fractionating a mixture of peptides eluted from a first MHC class II allotype;
    - (b) identifying a self peptide from said mixture;
  - (c) testing whether said self peptide binds to a second MHC class II allotype, said binding being an indication that said self peptide is a nonallelically restricted immunomodulating peptide.

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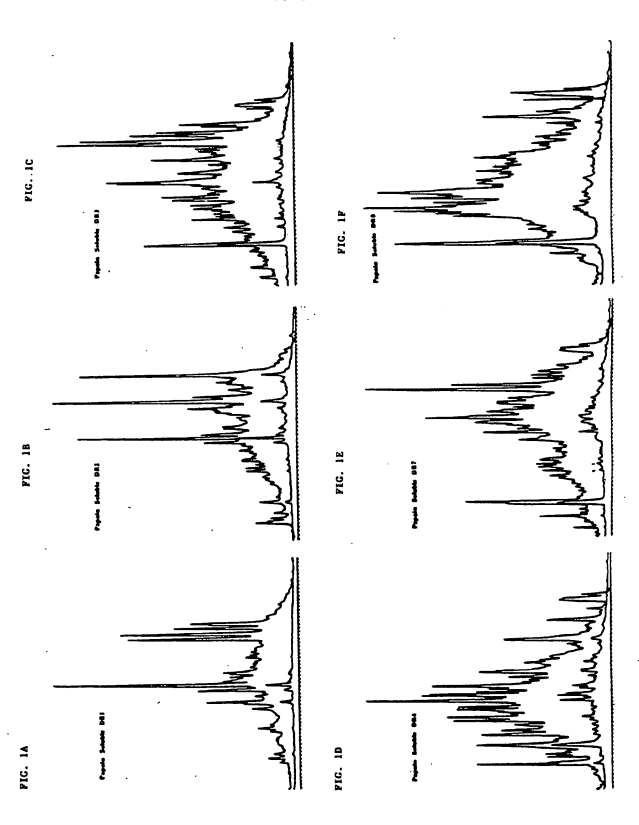
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- 19. A method of identifying a potential immunomodulating peptide, said method comprising:
- (a) providing a cell expressing MHC class II molecules on its surface;
- (b) introducing into said cell a nucleic acid encoding a candidate peptide;
- (c) determining whether the proportion of said class II molecules which are bound to said candidate peptide is increased in the presence of said nucleic acid compared to the proportion bound in the absence of said nucleic acid, said increase being an indication that said candidate peptide is a potential immunomodulating peptide.
- 20. A method of identifying a potential immunomodulating peptide, said method comprising:
- (a) providing a cell expressing MHC class II molecules on its surface;
- (b) introducing into said cell a nucleic acid encoding a candidate peptide;
- (c) determining whether the level of MHC class II molecules on the surface of said cell is decreased in the presence of said nucleic acid compared to the level of said molecules in the absence of said nucleic acid, said decrease being an indication that said candidate peptide is a potential immunomodulating peptide.
- 21. A method of identifying a nonallelically restricted immunostimulating peptide, said method comprising:
- (a) providing a cell bearing a first MHC class I or class II allotype, said cell being infected with a pathogen;
- (b) eluting a mixture of peptides bound to said cell's first MHC allotype;

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- (c) identifying a candidate peptide from said mixture, said candidate peptide being a fragment of a protein from said pathogen;
- (d) testing whether said candidate peptide binds to a second MHC allotype, said binding being an indication that said candidate peptide is a nonallelically restricted immunostimulating peptide.





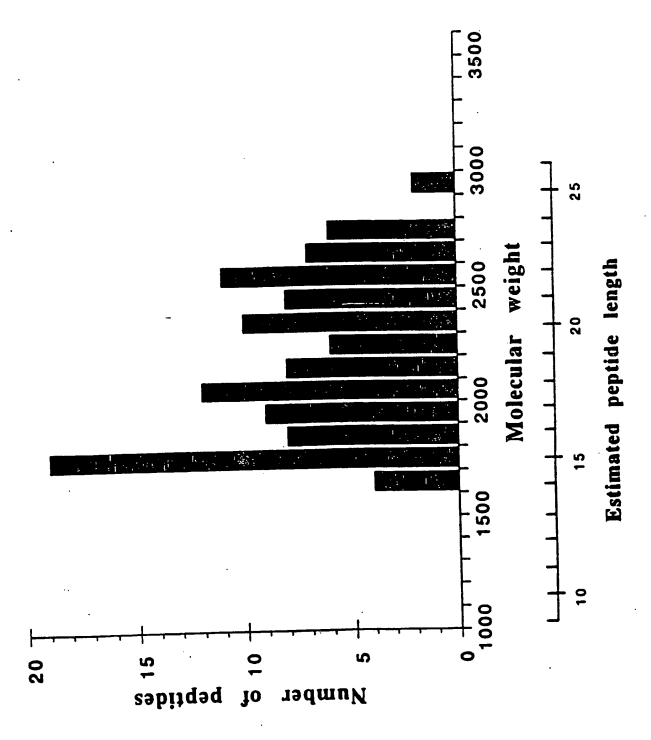


FIG. 2

3/3.

FIG. 3A

A.

ATG GCC ATA AGT GGA GTC CCT GTG CTA GGA TIT TTC ATC ATA GCT N A I S G V P V L G P F I I A

GTG CTG ATG AGC GCT CAG GAA TCA TGG GCT AAG ATG CGC ATG GCC V L N S A Q B S N A K H R H A

ACC CCG CTG CTG ATG CAG GCG CTG CCC ATG TAA

T P L L N Q A L P N STOP

FIG. 3B

B.

ATG GCC ATA AGT GGA GTC CCT GTG CTA GGA TTT TTC ATC ATA GCT M A I S G V P V L G P F I I A

GTG CTG ATG AGC GCT CAG GAA TCA TGG GCT CTT CCC AAG CCT CCC V L M S A Q B S W A L P K P P

AAG CCT GTG AGC AAG ATG CGC ATG GCC ACC CCG CTG CTG ATG CAG K P V S K H R H A T P L L M Q

GCG CTG CCC ATG TAA
A L P M stop

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# INTERNATIONAL SEARCH REPORT

E-mational application No.
PCT/US93/07545

IPC(5)	SIFICATION OF SUBJECT MATTER A61K 37/00, 37/02, 37/22, 31/70; C07K 7/00, 7/08, 7 530/300, 324, 325, 326, 327; 514/2, 44; 536/23.1; 424 International Patent Classification (IPC) or to both nat	I/450	,	
B. FIEL	DS SEARCHED			
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APS. Med	line, Biosis, Chem Abs, Derwent WPI, Embase, search on, author names, antigen presentation, autoimmune	terms: peptide, MHC, class II, DR, I-A	, I-E, liposome, iscom,	
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appr	opriate, of the relevant passages	Relevant to claim No.	
Y	Journal of Immunology, Volume 145, Number 6, Issued 15 September 1990, D.O. Sullivan et al., "Characterization of the specificity of peptide binding to four DR haplotypes", pages 1799-1808, see entire document.			
Y	Immunology Today, Volume 12, Number 11, issued November 1991, A.M. Mowat et al., "ISCOMS - a novel strategy for mucosal immunization?", pages 383-385, see entire document.			
Y	Immunology Today, Volume 11, issued al., "Peptide competition for antigen presentire document.	January 1990, L. Adorini et sentation", pages 21-24, see	1-18	
X Further documents are listed in the continuation of Box C. See patent family annex.				
T later decorment published after the international filing date or priority				
*A* document defining the general state of the art which is not considered principle or theory underlying the investion			rvestice	
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ategory*	Citation of document, with indication, where appropriate, of the relevant	ant passages	Relevant to claim No.
·	Journal of Immunology, Volume 148, issued 01 June Collins et al., "Processing of exogenous liposome encarantigens in vivo generates class I MHC-restricted T ce responses", pages 3336-3341, see entire document.	psulated	10
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